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UTILITY PATENT APPLICATION TRANSMITTAL

(Only for new nonprovisional applications under 37 C.F.R. § 1.53(b))

Attorney Docket No.	TB-104IB
First Inventor or Application Identifier	Jamie Grooms
Title	Cortical Bone Cervical Smith-Robinson Fusion Implant
Express Mail Label No.	EK670287616US

APPLICATION ELEMENTS

See MPEP chapter 600 concerning utility patent application contents

ADDRESS TO: Assistant Commissioner for Patents
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Washington, DC 20231

- ☒ * Fee Transmittal Form (e.g., PTO/SB/17)
(Submit an original and a duplicate for fee processing)
- ☒ Specification [Total Pages **31**]
(preferred arrangement set forth below)
 - Descriptive title of the Invention
 - Cross References to Related Applications
 - Statement Regarding Fed sponsored R & D
 - Reference to Microfiche Appendix
 - Background of the Invention
 - Brief Summary of the Invention
 - Brief Description of the Drawings (if filed)
 - Detailed Description
 - Claim(s)
 - Abstract of the Disclosure
- ☒ Drawing(s) (35 U.S.C. 113) [Total Sheets **22**]
- Oath or Declaration [Total Sheets **1**]
 - ☐ Newly executed (original or copy)
 - ☐ Copy from a prior application (37 C.F.R. § 1.63 (d))
(for continuation/divisional with Box 16 completed)
 - ☐ **DELETION OF INVENTOR(S)**
Signed statement attached deleting inventor(s) named in the prior application, see 37 C.F.R. §§ 1.63(d)(2) and 1.33(b).

- ☐ Microfiche Computer Program (Appendix)
- Nucleotide and/or Amino Acid Sequence Submission (if applicable, all necessary)
 - ☐ Computer Readable Copy
 - ☐ Paper Copy (identical to computer copy)
 - ☐ Statement verifying identity of above copies

ACCOMPANYING APPLICATION PARTS

- ☐ Assignment Papers (cover sheet & document(s))
- ☐ 37 C.F.R. § 3.73(b) Statement (when there is an assignee) ☐ Power of Attorney
- ☐ English Translation Document (if applicable)
- ☐ Information Disclosure Statement (IDS)/PTO-1449 ☐ Copies of IDS Citations
- ☐ Preliminary Amendment
- ☒ Return Receipt Postcard (MPEP 503)
(should be specifically itemized)
*Small Entity Statement(s) ☐ Statement filed in prior application, Status still proper and desired
- ☒ Statement(s) (PTO/SB/09-12)
- ☐ Certified Copy of Priority Document(s) (if foreign priority is claimed)
- Other:

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16. If a CONTINUING APPLICATION, check appropriate box, and supply the requisite information below and in a preliminary amendment:

Continuation Divisional ☒ Continuation-in-part (CIP) of prior application No: **15 98/17769 +**

Prior application information: Examiner

Group/Art Unit:

For CONTINUATION or DIVISIONAL APPS only: The entire disclosure of the prior application, from which an oath or declaration is supplied under Box 4b, is considered a part of the disclosure of the accompanying continuation or divisional application and is hereby incorporated by reference. The incorporation can only be relied upon when a portion has been inadvertently omitted from the submitted application parts.

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ADDITIONAL FEES</p> <table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th colspan="2">Large Entity</th> <th colspan="2">Small Entity</th> <th rowspan="2">Fee Description</th> <th rowspan="2">Fee Paid</th> </tr> <tr> <th>Fee Code</th> <th>Fee (\$)</th> <th>Fee Code</th> <th>Fee (\$)</th> </tr> </thead> <tbody> <tr> <td>105</td> <td>130</td> <td>205</td> <td>65</td> <td>Surcharge - late filing fee or oath</td> <td></td> </tr> <tr> <td>127</td> <td>50</td> <td>227</td> <td>25</td> <td>Surcharge - late provisional filing fee or cover sheet</td> <td></td> </tr> <tr> <td>139</td> <td>130</td> <td>139</td> <td>130</td> <td>Non-English specification</td> <td></td> </tr> <tr> <td>147</td> <td>2,520</td> <td>147</td> <td>2,520</td> <td>For filing a request for reexamination</td> <td></td> </tr> <tr> <td>112</td> <td>920*</td> <td>112</td> <td>920*</td> <td>Requesting publication of SIR prior to Examiner action</td> <td></td> </tr> <tr> <td>113</td> <td>1,840*</td> <td>113</td> <td>1,840*</td> <td>Requesting publication of SIR after Examiner action</td> <td></td> </tr> <tr> <td>115</td> <td>110</td> <td>215</td> <td>55</td> <td>Extension for reply within first month</td> <td></td> </tr> <tr> <td>116</td> <td>380</td> <td>216</td> <td>190</td> <td>Extension for reply within second month</td> <td></td> </tr> <tr> <td>117</td> <td>870</td> <td>217</td> <td>435</td> <td>Extension for reply within third month</td> <td></td> </tr> <tr> <td>118</td> <td>1,360</td> <td>218</td> <td>680</td> <td>Extension for reply within fourth month</td> <td></td> </tr> <tr> <td>128</td> <td>1,850</td> <td>228</td> <td>925</td> <td>Extension for reply within fifth month</td> <td></td> </tr> <tr> <td>119</td> <td>300</td> <td>219</td> <td>150</td> <td>Notice of Appeal</td> <td></td> </tr> <tr> <td>120</td> <td>300</td> <td>220</td> <td>150</td> <td>Filing a brief in support of an appeal</td> <td></td> </tr> <tr> <td>121</td> <td>260</td> <td>221</td> <td>130</td> <td>Request for oral hearing</td> <td></td> </tr> <tr> <td>138</td> <td>1,510</td> <td>138</td> <td>1,510</td> <td>Petition to institute a public use proceeding</td> <td></td> </tr> <tr> <td>140</td> <td>110</td> <td>240</td> <td>55</td> <td>Petition to revive - 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Signature				Date

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1.0 Background of the Invention

U.S. Patent No. 5,306,309, discloses a spinal disk implant comprising a solid body of biocompatible synthetic material arranged to define a right-rectangular solid having two opposed side faces and two opposed transverse faces, including a convexly curved anterior face and a

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posterior face, for implantation in the intervertebral space. The discussion of vertebral and intervertebral morphology is hereby incorporated by reference.

U.S. Patent No. 5,609,635, discloses a lordotic interbody spinal fusion implant comprising a wedge shaped metallic cage for insertion into the intervertebral space.

U.S. Patent No. 5,306,307, discloses a ceramic spinal disk implant having a serrated edge.

None of these references disclose a cortical bone intervertebral implant having a substantially D - or bread-loaf-shaped structure having a canal into which osteogenic, osteoinductive, or osteoconductive materials may be packed, which sustains spinal loads, and which is remodeled into the spine in the course of fusion. Accordingly, the present invention addresses the need in the art for improvements to both the implant and the avoidance of post-surgical complications from anterior cervical fusions. The present invention provides a new cortical bone implant for use in achieving anterior cervical fusions when implanted according to the Smith-Robinson procedure. In addition, in view of the peculiar characteristics of bone, the present invention comprises unique methods and apparatuses for the manufacture of the substantially "D"-shaped cortical bone implant.

2.0 Summary of the Invention

An implant composed substantially of cortical bone is provided for use in cervical Smith-Robinson vertebral fusion procedures. According to methods of this invention, the implant is derived from allograft or autograft cortical bone sources, is machined to form a substantially D - or other appropriately shaped implant having a canal into which osteogenic, osteoinductive, or osteoconductive material may be packed. The implant is inserted into the space between adjacent cervical vertebrae to provide support and induce fusion of the adjacent vertebrae.

3.0 Brief Description of the Drawings

Figure 1 provides several views of the fusion implant of this invention.

Figure 2 provides views of the core cutter and drill assembly and the bone plug formed by cutting into the diaphysis of a long bone when such a core cutter and drill assembly is used.

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Figure 3 provides a view of broach as used according to this invention and an asymmetric canal formed by use of such a broach.

Figure 4 provides several views of an apparatus for machining a profile on the exterior surface of the implant of this invention.

5 Figure 5 provides a view of an apparatus for inscribing retention teeth in the upper surface, lower surface or both upper and lower surfaces of the implant.

Figure 6 provides several views and dimensions for specific embodiments of the implant of this invention.

Figure 7 provides a view of a stacked embodiment of the implant of this invention.

10 Figure 8 provides several views of an implant of this invention formed by juxtaposition of mirror image halves of the implant, as well as an embodiment useful for posterior lumbar intervertebral fusion procedures (PLIFs).

Figure 9 provides a view of a stacked embodiment of the implant of this invention wherein the stacked constituents thereof are retained in registered relationship by press-fitting or otherwise
15 bringing more than one implant into contact with each other and having a cancellous plug or other biocompatible material located in the central canal of each stacked implant, thereby acting as a retention pin.

Figure 10 shows an alternate method for producing bone stock for making the implant of this invention of essentially unlimited height from the anterior margin of the tibia or the linea aspera of
20 the femur.

Figure 11 shows dimensions and further processing of the implant of this invention produced according to the alternate method of figure 10.

Figures 12-17 show final profiles for implants produced according to the alternate method of figures 10 and 11.

25

4.0 Detailed Description of the Invention

According to this invention, a substantially "D"-shaped cortical bone implant for cervical Smith-Robinson fusions is produced, preferably under aseptic conditions. Class 10 clean room
30 processing is desirable, and sterilization of all machining tools is likewise preferred, (particularly

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after switching from one allograft donor to the next), so that the finished product may be treated by standard techniques known in the art (alcohol, peroxide, or like treatments), prior to storage and shipment to physicians for use in implantation procedures. Because of the peculiarities of working with bone, and in particular, because of the desirability of maintaining aseptic conditions while
5 working with this material, novel approaches have been adopted in the production of the product of this invention.

The implant is preferably formed from cortical bone obtained from tibia, femur or other source of strong cortical bone. The bone source may be autograft or, due to possible complications at the donor site (infection, pain, delayed healing), is preferably, allograft bone. In addition, it is
10 critical that the source bone be derived from a donor whose medical history is well known (absence of transmissible diseases, cancer, osteoporosis), and that the donor bone be obtained under aseptic conditions according to accepted practices in the art of tissue banking. In addition, extensive *in vitro* testing should be conducted to ensure the absence of pathogenic agents.

The approach adopted in describing the implant of this invention is to first provide a
15 narrative disclosure of preferred methods for making the implant, followed by a detailed description of the implant itself, followed by a detailed description of various apparatuses and aspects of the machining process, and finally, a detailed description of the method of using the implant.

4.1 Narrative Description of Implant Manufacture

20 While any shape of cortical bone may be used to begin with, we have found that for consistent production of cortical bone which may be reliably machined, it is advantageous to commence with a plug of bone which extends from the exterior of the diaphysis of a long bone toward the intramedullary canal (where, *in vivo*, the bone marrow resides). The result is a bone plug
25 or dowel which has an outer substantially cortical end and an internal end which is composed largely of soft cancellous bone. In cutting the bone plug, we have discovered that the use of a core cutter is convenient. This device comprises an outer coring element of any desired diameter, whereby the diameter of the bone plug is defined, and a centrally located solid drill bit, which provides a canal through the center of the bone plug as well as stability for the core cutting element.

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The core cutter-drill assembly is preferably torqued by an air drill, driven by sterile air, and the source bone is preferably immobilized in a sterilized vice during the core-cutting process.

We have discovered that in the above-described manner, cortical bone implants may be fashioned having heights, widths and lengths which are practically useful in the Smith-Robinson cervical fusion method. According to this method, the height of the implant is only limited by the distance from the exterior of the bone diaphysis to the intramedullary canal. However, we have discovered that, by this method, final implant heights from about 7 mm to about 14 mm may be produced, depending on the choice of bone source and the location on the bone from which the bone plug is cored. Since it is extremely rare for the cervical intervertebral space to extend beyond these limits, this method is therefore capable of supplying implants of required or useful heights.

Likewise, the length and width of the implant are defined by the diameter of the core-cutter, and final lengths and widths of between about 7 and 14 mm are easily provided for by this method. In addition, where the need arises for heights between about 10 mm and 14 mm, or if difficulty is experienced in obtaining donor bone having a sufficient width from the exterior of the bone to the intra-medullary canal to provide such heights, alternate methods of producing the implant of desired heights disclosed herein may be employed. For example, in a first such alternate method, implants of this invention are produced and then stacked to provide a unitary implant of the desired height dimensions. Such stacked implants may be maintained in a unitary association by drilling appropriate holes through the height of the implant, and inserting therein appropriate retention pins made from any desirable material, including cortical bone, bioabsorbable synthetic polymer, titanium or other metallic retention pins. Alternatively, the stacked implants may be retained in a unitary association by means of a plug of cancellous bone, hydroxyapatite or other biocompatible, osteoconductive or osteoinductive material, and press-fitting the stacked implants to achieve the desired height (see figure 9). In a further alternate method, a section of cortical bone along the long axis of a long bone may be machined according to methods known in the art. By then further shaping and cutting appropriate heights in such cortical bone, and bringing halves of the implant into juxtaposition with each other, implants of any desired shape and height are produced. In yet a further alternate procedure, (see figures 10-17), unitary implants of this invention of essentially unlimited height are produced by length-wise sectioning the anterior margin of the tibia or linea aspera of the femur, segmenting the substantially triangular cortical bone to desired heights, drilling

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a cannulation through the segments thus produced, and finally shaping the implants to desired dimensions, as defined below for the first principal method of making the implant of this invention.

Continuing with a description of the first method for making the implant of this invention, the cancellous bone on the internal side of the bone plug is removed by any convenient means, including with a saw, an abrasive means such as a diamond tipped rotary sander, or a tooling bit mounted in a lathe, to produce a "washer" shaped piece of substantially cortical bone. Both the internal and external ends of the bone plug should be machined flat, thereby forming a top face and a bottom face, each of which is substantially planar, and preferably parallel. While the cancellous bone is partially or completely removed by this process, there remains a slight difference in the density of the bone from the external (cortical) to the internal (cancellous or originally intra-medullary) aspect of the bone plug. It is desirable to record the orientation of the bone plug as subsequent machining steps proceed most efficiently when machined from the external aspect toward the internal aspect.

In order to accommodate subsequent machining steps and to provide an orientation to the implant according to which the surgeon may properly insert the implant, the circular internal canal formed by the centrally located solid drill bit of the core-cutter is modified to form an asymmetric shape, such as a key way. This may be achieved by any of a number of different means, including drilling a slot into an aspect of the internal canal closest to the external (more dense cortical) end of the dowel. In one embodiment of this invention, we have found that an implant of consistently good final quality may be machined by conversion of the circular canal into a substantially "D" shaped canal having three essentially rectangular walls and a fourth convexly curved wall. We have found that it is desirable for the curvature of the convexly curved wall to approximate the external curvature of the bone plug. This modification may be achieved by any of a variety of means. However, we have invented an efficient means by which consistently usable implants may be reproducibly machined. This is accomplished by immobilizing the implant, for example in an arbor press assembly, and, preferably from the originally cortical external (denser) end of the implant, slowly forcing a broach through the originally circular canal. The broach is preferably a hard metallic member having a plurality of spaced-apart ribs or rings machined therein, with indentations provided between each ring which thereby form the spacing between adjacent rings. In addition, the edges of each ring are desirably very precise, angular, and sharp, such that as the broach is forced

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through the originally circular internal canal, the sharp cutting edge of each ring shaves off an incremental amount of bone as the ring passes through the implant. Each ring of the plurality of rings has a shape which, starting at the insertion end of the broach is tapered from an essentially circular shape to any desired final shape for the canal. Accordingly, in one embodiment of this invention, the rings transition from a circular shape to a substantially "D"-shaped profile or any other desired shape over several inches and over a plurality of spaced apart rings. It will be appreciated that the length of the broach and the number of rings used is defined by the amount of bone that must be removed to form the new shape, the width of each ring and the width of the space between each ring. Removal of no more than about 0.004" of bone by each ring has been found to be a sufficiently small transition to ensure that the vast majority of implant blanks survive this machining step. Broaches of approximately 6" in length have been found adequate for most implant shapes, but for very asymmetric shapes (e.g. an implant which is 11 mm wide and 14 mm long), more bone would need to be removed to form the "D"-shaped canal than from a symmetric implant (e.g. a 14 mm wide by 14 mm long implant). This need may be accommodated by use of more than one broach, with the shape of the insertion end of each consecutive broach substantially matching the shape of the last ring on the previous broach.

Having formed an asymmetric shape, such as a key way, from the internal canal running through the implant, we have found it desirable to modify the external profile of the implant from a substantially circular shape to another desired form. In one embodiment of this invention, the external form of the implant is machined so as to proportionately match the shape of the substantially "D"-shaped internal canal. An external "D"-shaped profile has been used in implants known in the art (see for example U.S. patent 5,306,309; 5,522,899) made from materials other than bone, because of the ability of the convexly curved face of the implant to substantially match the curvature of the anterior aspect of the intervertebral disk into which the implant is to be inserted, as well as to provide efficient spinal load distribution over the remainder of the implant. However, due to the peculiar nature of bone, and the requirements of aseptic or sterile manufacturing, inventive methods and apparatuses were required to produce the desired external profile for the cortical bone implant. It will be recognized that, based on the instant disclosure, a substantially "D"-shaped external profile of the implant may be machined by a variety of means which vary from the precise methods disclosed herein. In addition, other external profiles than the "D"-shaped profile are

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likewise enabled by modifications of the methods and apparatuses disclosed herein for formation of the "D"-shaped external or internal profile. Thus, according to one alternate method of making the implant of this invention, for example where the linea aspera of the femur is sectioned or where the anterior margin of the tibia is sectioned, an external profile that substantially varies from a "D"-
5 shaped device may be produced, (see figures 10-17).

We have found it convenient and reproducible to use either of two principal methods for machining the external profile. The implant, with the "D" or alternately shaped internal canal being used as a key way, is fitted onto the end of a spindle which precisely matches the shape of the internal canal of the implant, thereby providing purchase for machining of the external profile of the
10 implant. In a first preferred method, as the implant is rotated on the spindle, it is contacted with an asymmetric generator (grinding) wheel attached to a cog which meshes at a known registration point with a cog to which the spindle with the implant is attached. The speed of rotation of the exterior of the spindle mounted implant, and the exterior of the generator wheels are designed to differ such that as the generator wheel and implant are contacted and are rotated in fixed registration, the
15 generator surface (which is preferably an abrasive diamond plated surface), grinds bone from the external surface of the implant, to form a profile thereon defined by the asymmetric shape of the grinder wheel.

In a second external profile generation method, the implant, with the "D" or alternately shaped internal canal being used as a key way, is fitted onto the end of a spindle which precisely
20 matches the shape of the internal canal of the implant, thereby providing purchase for machining of the external profile of the implant. In this method, the spindle is affixed to an asymmetric cam which rotates concentrically with the spindle, and therefore the implant. The thus mounted implant is contacted with a cutter means, such as a sharp bit having cutting edges which rotate about an adjacent axis. The implant mounted spindle riding on the asymmetric cam is biased to contact the
25 rotating cutter, which thus traces a profile onto the exterior of the implant defined by the shape of the asymmetric cam. For purposes of this disclosure, use of the term "asymmetric cam" should be understood to mean any desirable shape such that upon production of the implant, the shape thereof is defined by that of the asymmetric cam. Shapes contemplated by this disclosure include, but are not limited to, elliptical shapes, D-shapes, partially curved shapes, and the like. The implants
30 produced according to any of the alternate procedures are likewise shaped, although the final shape

may vary depending on the size of the bone stock used (see for example the final shapes of the device shown in figures 12-17).

Once the external profile has been machined, the implant is removed from the spindle, and the machining of the implant may either be terminated, to provide a substantially "D"-shaped cortical bone implant with flat upper and lower surfaces, or an external feature may be machined into the upper and lower surfaces to prevent backing out of the implant upon insertion into the intervertebral space. This may be achieved by a number of means, such as by machining annular rings, indentations and projections, ribbing or teeth into the upper, lower, or both surfaces of the implant. In a preferred embodiment of this invention, the implant is passed through a set of opposing jaws bearing teeth which broach a tooth-shaped profile into the implant as it is forced through the jaws. Alternatively, the implant is passed several times over a ridged surface which cuts the desired tooth profile into the upper, lower or both surfaces of the implant. Preferably, the thus formed teeth angle toward the anterior (convexly curved) face of the implant to prevent backing out of the implant once it is inserted into an appropriately shaped cavity formed in the intervertebral space in an anterior aspect of the cervical spine. In order to accommodate the difficulty surgeons experience in forming precise angles when forming such cavities in the spine, (see for example U.S. patent No. 5,397,364 disclosing a beveled edge to reduce trauma upon insertion of a metallic spinal implant), a beveled edge of defined radius is preferably machined into three faces of the implant, but leaving the anterior face unbeveled. The sharp anterior edge, like the teeth in the upper and lower surfaces of the implant, retards backing out of the implant.

4.2 Detailed Description of the Implant

Referring now to figure 1A, there is shown a top view, as if viewed from the top of the spinal column, of a substantially D-shaped cortical bone implant 100. The implant has a wall thickness 101, a length 103, a width 102, and an internal canal 104, which fall within desired tolerances (see discussion below). The implant comprises four contiguous walls, including a substantially straight rear wall 105, substantially straight side walls 106 and 107, and a preferably curved front wall 108. In figure 1B, there is shown a side view of the implant 100, revealing the height 109, of the implant. In addition, this view shows, in outline, the internal side walls 106' and 107' of the internal canal,

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104. It also shows the top **110** and bottom **111** surfaces of the implant. In figure 1C, there is shown a top view of an embodiment of the implant **100** in which an external feature **120** has been inscribed onto the top **110** and bottom **111** surfaces of the implant. In addition, a "radius" or bevel **115** is shown on the two side and posterior edges of the implant. Figure 1D shows a side view of the
5 implant **100** in which the inscribed feature **120** can clearly be seen in the top **110** and bottom **111** surfaces of the implant. In this view, it can be seen that the external feature **120** has the side profile of a set of teeth, all of which angle toward the anterior face **108** of the implant. An outline of the bevel **115** is also evident in this view, as is the rounded posterior edge **105**. As can be seen, in this embodiment of the invention, the anterior edge **108** is maintained with a sharp edged. In figure 1E,
10 there is shown a detail of one embodiment of the inscribed feature **120** on the portion of the implant indicated in figure 1D. In a preferred embodiment, the feature **120** defines a tooth-like structure, with teeth **121** separated from each other by concavities **122**. An angle defines the grade of the concavity as it ramps to the tooth. The tooth height **123**, space between teeth **124**, and aperture of the concavity **125** may all be defined by the manufacturer to optimize retention of the implant within
15 the cervical spine after proper placement.

4.3 Detailed Description of the Method of Manufacturing the Implant

Because of the peculiar nature of bone, and the desirability of sterile or aseptic
20 manufacturing, specific and specialized procedures and apparatuses are required for successful formation of the implants of this invention. Those skilled in the art will recognize that, based on the methods and apparatuses disclosed herein, the implant of this invention may be manufactured by alternate means suggested by those described herein. Nonetheless, through careful design and knowledge of bone structure, instruments for the manufacture of the implant of this invention have
25 been invented for this purpose. In what follows, specific details with respect to preferred method and apparatuses for making the implant of this invention are provided. It should be recognized that the invention should not be construed as being limited to these specifics.

Referring to figure 2, there is shown in side view in figure 2A a core cutter **200**, having a core bit **201** which is affixed by a set screw **203** to the shaft **204** of a drill bit **202**, centrally located
30 within and coaxial with the core cutter. In figure 2B, an end-on view of the core cutter **200** is

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provided showing the set screw **203** in outline. Figure 2C shows a side view of the bone plug **210** which is formed by cutting a plug of bone from the diaphysis of a long bone using the core cutter **200**. At one end, **211**, originally the external cortical surface of the bone shaft, there is a substantially cortical bone surface through which a hole **213** is formed by the central bit **202** of the core cutter **200**. The other end, **212**, is an irregular and bone surface which, *in vivo*, formed part of the wall of the intramedullary canal. Cancellous bone or other microstructure at the end **212** is removed, and both ends are ground, cut or otherwise machined to be substantially flat and parallel, to form the substantially cortical bone plug **210** shown in figure 2D.

Referring to figure 3, there is shown in figure 3A an internal canal profile broaching tool **300**. A plurality of spaced-apart ribs or rings **301** are provided along the length of the broach which taper from a substantially circular shape at the insertion end **302** of the broach, to substantially "D"-shaped rings **303** (or any other desired shape) at the completion end **304** of the broach **300** (intermediate ribs **305** are not shown; rather, the outline of the taper angle is shown). A notch or groove **306** is provided in the broach completion end **304** for releasably affixing the broach into a means, such as a press, for forcing the broach through the implant canal. In figure 3B, there is provided an end-on view of the cancellous bone plug **310** after the broaching procedure is completed. As can be seen, the internal canal **104** has been converted from a circular canal into a substantially "D"-shaped canal. As will be appreciated from this disclosure, any of a number of different asymmetric shapes in the internal canal **104** may be defined by this or analogous means, the principal goal being to provide a purchase (referred to herein as a "key way") within the implant for external machining of the implant. In one embodiment (see figure 9), the canal may be retained as a substantially circular canal, and a slot **904** is machined therein to provide the necessary asymmetry to form a key way.

Having formed a key way within the implant, it is possible to modify the external profile of the implant. In one aspect of this invention, referring to figure 4A, this is conveniently achieved by affixing the implant **410** to the spindle **420** of a lathe **400**. The spindle shaft **440** extends, through bearings (not shown), to a means **450** (such as a handle or a motor) for rotating the spindle. Affixed to the spindle-shaft is a cam **430**, the shape of which defines the ultimate external profile of the implant **410**. The spindle shaft **440** and bearings are mounted in a cross slide **441**, which translates in a first plane, referred to as the "Y-plane". Motion in the Y-plane is limited by contact of the cam

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430 with a limiting means **460** such as a cam follower, which remains in register with a carriage **442** which translates along a plane, the "X-plane", transverse to the Y-plane motion of the cross-slide . The cross-slide is mounted in a slide-way **443** of the carriage **442**, which in turn is slideably mounted on the bed **444** of the lathe, such that the carriage **442** is permitted to translate along the X-plane. Travel of the slide **442** along the X-plane is limited by means of a stop screw **470**.

Further detail of this means for generating the external profile of the implant is provided in figure 4B, which provides a side view of one specific embodiment of the implant external profile generator **400**. An air driven turbine within housing **401** provides a source of torque to turn a shaft **402**. A means for cutting or grinding the external surface of the implant **410**, such as an appropriately fashioned cutter or bit having a non-cutting end **403** for fixation to the shaft **402**. Extending from the non-cutting end **403** which has a first diameter, is a cutting surface **404**, having a second, smaller diameter. A "shoulder" **405**, forms a radius extending between the smaller diameter of the cutting surface **404** and the larger diameter of non-cutting surface **403**. The cutting surface **404** is contacted with the implant blank **410**, mounted on spindle **420**, to which, as described above is mounted an asymmetric cam **430**. The thus mounted implant blank **410** is brought into contact with the cutting surface **404**, by virtue of translation in the X-plane of the carriage **442**. The spindle **420**, and thus the asymmetric cam **430** are rotated, manually or by motor driven means, through shaft **440** and handle **450** which are attached concentrically with the cam **430**. Preferably, the asymmetric cam **430** is elastically biased toward a stationary cam follower **460**. In this fashion, after several revolutions of the handle **450**, the shape of the asymmetric cam **430** generates the desired external profile of the implant **410** riding on the spindle **420**, through contact with the rotating cutting surface **404**.

To ensure that the implant blank is machined only up to the point that the forward edge **411** of the implant approaches but does not contact the "shoulder" **405** on the cutter, a stop screw **470** is provided. The stop screw **470** is set to prevent further advancement of the implant blank **410** by stopping advancement of the carriage **442** when the leading edge **471** of the stop screw comes into contact with a measuring screw **480**. The appropriate setting of the stop screw **470** is achieved at the start of the milling process by first placing the implant **410** between the end **481** of the measuring screw **480** and an anvil **482**, and tightening the measuring screw **480** until it just makes contact with the implant. In this fashion, the measuring screw **480** and anvil **482** essentially form a micrometer,

with the gap being defined by the width of the implant. Both the measuring screw **480** and anvil **482** are housed within a measuring slide **483** which, when slid all the way to the left as shown in figure 4B, abuts a rotateable stop cam **490**, retained within the same slide-way as the measuring slide **483** by a retainer **484**. The rotateable stop cam **490** may be set in either of two positions,

5 which produces a difference in the stopping point of the stop screw **470** of approximately 0.06".

The significance of this difference is that the first position arrests advancement of the stop screw **470** (and therefore the carriage **442**) just before the implant **410** contacts the radius shoulder **405** of the cutting surface **404**. In the second position, the stop cam **490** allows the stop screw to advance the additional approximately 0.06" to allow contact of the implant **410** with the shoulder **405** of the cutting surface **404** to thereby bevel the edges of the implant **410** that are thus contacted.

10 Accordingly, in the pre-milling setup, the stop cam **490** should be rotated such that the stop screw **470** is forced to stop the extra 0.06", following which a further processing step may be carried out in which the stop cam **490** is rotated to the second position in which the stop screw **470** is allowed to advance this additional approximately 0.06".

15 In figure 4C, there is provided an end-on, rear view (i.e. looking from the handle **450** toward the spindle **420**) of the asymmetric cam **430**, the spindle **420** and the implant **410**. In addition, in this detail view, an additional feature in the asymmetric cam **430** is seen as a diminution in the thickness along three faces **431** of the asymmetric cam **430** which is a relief in the rear of the asymmetric cam **430**. The significance of this relief **431** is that it restricts the contact of the implant **410** with the shoulder **405** to the extent defined by the relief in the rear of the asymmetric cam **430**.

20 As noted above, in fashioning an implant site in the intervertebral space during a partial discectomy, surgeons are unable to produce perfectly sharp angles. To accommodate this imperfection, to prevent trauma upon insertion of an implant with sharp edges, and to create as tight-fitting an implant as possible, the fashioning of a bevel around the edges of the implant that are inserted into the intervertebral space created by the surgeon is desired. At the same time, in order to prevent backing out of the implant, it may be desirable to retain a sharp anterior implant edge, and therefore the relief in the cam **430** does not extend completely around the cam. Thus, upon completion of the external profile of the implant **410** as described above, the carriage **442** is backed away from the cutter, the stop cam **490** is flipped to its second position allowing advancement of the stop screw

30 **470** the additional approximately 0.06" mentioned above. At the same time, a shot pin **432** is

advanced into the relief **431** by means of a shot pin mover **433**, thereby allowing rotation of the cam **430** only to the extent permitted by the shot pin **432** as it rides within the relief **431**. With the shot pin **432** riding in the relief **431**, the "shoulder" **405** contacts the leading edge **411** of the implant blank **410**, thereby rounding three edges of the implant **410**. After machining the leading edge **411** of the implant **410**, the implant is removed from the spindle **420**, turned around, and re-positioned on the spindle **420**, to inscribe the bevel on three edges of the other side of the implant.

In figure 4D, a frontal view is provided of the spindle **420**, the implant **410**, the asymmetric cam **430**, and the cam follower **460**. Also shown is the cam adapter **461**, by means of which the cam follower **460** is affixed to the carriage **442**, and by means of which the cam follower **460** maintains the cutting surfaces **404/405** in contact with the implant **410** as defined by the shape of the asymmetric cam **430**. Also shown is a part of the cross-slide **441**, which is preferably biased or which may be pushed manually toward the cam follower **460**.

In figure 4E, a side detail view is provided of the stop cam **490**. In this view, a stop cam handle **491** is shown which allows the operator of the implant outside profile generator to fix the stop cam **490** in a first position **A**, and a second position **B**, whereby additional travel of the stop screw **470**, and thereby advancement of the carriage **442**, is provided in position **B**, of about 0.06" due to the difference in the distances shown for these positions.

By means of the apparatuses and method described above, a cortical bone implant **100** as shown in figure 1 having a substantially "D"-shaped external profile, and a substantially "D"-shaped internal canal is produced. Naturally, based on this disclosure, those skilled in the art will appreciate that other shapes, both for the external profile and internal canal of the implant may be produced. For example, an ellipsoid is produced by the above described methods simply by modification of the shape of the asymmetrically shaped cam **430**, and the internal canal shape may be modified by drilling, routing, or broaching using a broach that tapers to any desired shape. The thus formed implant may be used after machining as described, followed by appropriate cleaning methods known in the art (e.g. bathing in alcohol, peroxide treatment etc.). In addition, however, it may be desirable to inscribe an external feature on the upper surface **110**, the lower surface **111**, or both. Such a feature may take any desirable form, such as annular rings, indentations, projections, ribbing or teeth. In a preferred embodiment, teeth sloping toward the anterior aspect **108** of the implant are inscribed onto the top **110** and bottom **111** surfaces of the implant by forcing the implant through

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opposed broaches bearing inscribing teeth. Alternatively, the upper **110**, lower **111** or both surfaces in turn may be repeatedly run, manually or by a machine-driven means, over an appropriately fashioned jaw bearing abrasive teeth such that the required profile of teeth are inscribed into the surfaces of the implant. Desirably, the successive teeth of the jaw are incrementally raised in height such that each tooth is only required to remove a small amount of bone (about 0.004" per tooth, to a total depth of 0.015"). In addition, it is preferred that the rake (angle of the teeth) be sufficiently sharp as to allow the implant to bite into the implantation site, without at the same time being so sharp as to be excessively brittle.

In figure 5, figure 5A, there is provided a top view of one side of one embodiment of blades **502** for use in a broach assembly **500** for inscribing teeth into the top **110**, bottom **111** or both surfaces of the implant. In outline, there is shown a lock-down handle **501** for clamping the assembly of blades **502** to a base **503**. By bringing a mirror image jaw into register with the depicted broach, a space is formed between the opposing teeth **502** at a distance sufficient to accommodate passage of the implant therebetween, provided that the teeth abrade recesses into the top and bottom surfaces of the implant **100**. To ensure proper engagement of the blades **502** and the implant **100**, there is provided a non-cutting surface **506** for contacting the implant **100** as it is introduced into the broach assembly **500**. The non-cutting surface **506** acts as a type of micrometer, forcing the cutting surfaces of the teeth **502** sufficiently apart to properly engage the implant as it passes through the broach assembly **500**. In figure 5B, there is provided a side view of an implant mounting device **504** having a "D"-shaped cavity **505** into which a "D"-shaped implant may be fitted for passage through the opposing jaws of the broaching jaw apparatus **500**. The resultant implant has the profile shown in figures 1C-1E.

In figures 5C-5E, there is shown an alternate apparatus and method for fashioning the retention teeth in the implant. In figure 5C, there is shown a carriage **510** having an appropriately dimensioned slot **520** for receiving the implant to be grooved. A tensioning screw **530** brings a retention arm **531** into juxtaposition with carriage housing member **532**, thereby clamping the implant into position within slot **520**. Through carriage housing members **532** and **533**, there is aligned a guide-rod **534** for guiding the carriage containing the implant as it is raked across a blade assembly **540**, over which said carriage **510** is made to pass. Said guide rod **540** also conveniently acts as a hinge, allowing the carriage **510** to swing upward for implant loading and also permitting

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the carriage to move down toward the base as the implant surface is cut on each successive pass of the carriage over said blade assembly 540. The blade assembly 540 is bolted within a base 550 over which said carriage 510 slides. Said base 550 also acts to receive fixation screws 551 and 552 which retain said guide rod 534 in place. A plurality of individual blades 560 are placed in a recess 554 in the base 550 and are maintained in registered position by retention screws 552 passing through retention holes 553 in each blade. Each blade 560 has an initial non-cutting surface 561, which is approximately 0.015" below the cutting surface 562, which in combination with said plurality of blades, forms a flat loading area for implant insertion into said slot 520. Figure 5D provides a side view of one blade 560, while figure 5E provides an end on view of the carriage 510 as it sits above the base 550. Accordingly, the implant is inserted into the slot 520 with the carriage 510 swung up from the base 550. The carriage is then swung down into the starting position with the implant making contact with the non-cutting surfaces 561 of the plurality of blades. The implant is depressed so that it is forced snugly against the non-cutting surface, and then tensioned into place with the retention screw 530. Thereafter, the carriage is slid several times over the base 550 such that the cutting surfaces 562 of the plurality of blades thereby inscribe the desired tooth structure into the top surface, the bottom surfaces or both (after switching the implant around) surfaces of the implant. When the metallic bottom of the carriage comes into contact with the base, the machining of the implant is complete.

In figure 6A-I, there is provided a view of three different cortical bone implants according to this invention having particular geometries by way of example and not limitation. In figure 6A, there is shown an example of an implant 600 which has a height of 7 mm, a width of 11 mm, and a length of 14 mm. In addition, dimensions of various radii of the implant are provided. Note the effect of the "shoulder" 405 of the cutter which produces the a 0.059" radius and indent profile 610 starting at the approximate center of the part and proceeding around to the opposite side of the implant, i.e. around three faces of the implant. In figure 6B, the implant 600 is shown as a side view, and in figure 6C, there is shown a detail view of the teeth. Identical descriptions apply to the 7mmX11mmx11mm views of the implants of figures 6D-6F and the 7mmX14mmx14mm implant of figures 6G-6I.

In figure 7, there is shown a further aspect of this invention in which an implant, either machined as described above, or prior to said machining, is further machined so as to allow stacking

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thereof to achieve implants of various heights. Commencing from a blank cortical plug at the stage shown in figure 2D has the advantage that if breakage of the implant occurs during machining, this will likely occur prior to completion of all of machining steps. According to this embodiment of the invention, two implant blanks of known height are selected such that a unitary implant composed of both starting implants can be produced of a new desired height (e.g. a 6 mm high implant may be stacked with a 7 mm high implant to produce a 13 mm implant). Each implant blank is placed in a drill jig, and by means of a drill press or like means, holes are drilled through the implants. With the implants still in the jig, the jig is placed on the table of an arbor press. Pins, composed of cortical bone, resorbable but strong biocompatible synthetic material, or metallic pins of the appropriate diameter are then impelled into the holes in the implants such that the implants are formed into a unitary body by these pins. In order to encourage bony ingrowth, channels may be cut into the adjacent surfaces of the implants. The embodiment shown in figure 7A is a top view of an implant 700 into which four holes 701-704 have been drilled. In figure 7B, there is shown the juxtaposition of two implants 700A and 700B, with the drilled holes 701-704 in register to receive pins for maintaining the implants in register. In this view, the adjacent surfaces 710A and 710B have not been inscribed with teeth, while the surfaces 711A and 711B have been so inscribed. Based on this disclosure, those skilled in the art will recognize that a number of variations and modifications may be made to stack various forms of bone implants, or to maintain such implants in register with each other. These modifications are to be considered within the scope of this invention. Thus, as shown in figure 9, an implant 900 is produced by producing two implants 901 and 902, each having a cavity or canal 903, including an asymmetric key way 904 machined therein. By press-fitting the two implants together using an appropriately shaped cancellous plug 905 or a plug made from another biocompatible material, including but not limited to hydroxyapatite, cortical bone, synthetic materials, ceramic, optionally treated with growth factors such as bone morphogenetic protein and the like, the two implants 901 and 902 are retained in registered juxtaposition to form the implant 900.

In a further embodiment of this invention, shown in figure 8, a method for assembling the implant of this invention from component parts is provided. In figure 8A, there is shown an implant 800 composed of two side-by-side halves, 801A and 801B. The two halves of the implant are brought into juxtaposition to form a unitary implant. The two halves may be implanted in

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juxtaposition, or holes may be formed in each half, and the halves maintained in contact by forcing pins through the holes, in a fashion analogous to that described above for maintaining stacked implants in contact with each other. For this embodiment of the invention, a portion of cortical bone may be harvested from any suitable source of cortical bone. As shown in figure 8B, a segment, in the form of a block or a column of cortical bone is harvested along the long axis of a long bone, such as the femur, tibia, or fibula. The shape of the bone may be inscribed into the thus-harvested cortical bone by routing, broaching or other means as described herein. The thus-machined cortical bone may then be sectioned into appropriate heights, as needed, to provide the implant halves **801A** and **801B**. Alternate sites for harvesting the cortical bone segment are shown in figures 8B and 8C.

In addition to use for cervical Smith-Robinson type fusion, implants comprising each element, **801A** or **801B** alone, modifications and variations thereof, optionally in combination with another vertebral fusion implant, may be implanted, for example, to assist in induction of posterior lumbar intervertebral fusion (PLIF). In such a case, a device **810**, such as that shown in figure 8D-8G is machined from bone stock as shown in figures 8B, 8C or another appropriate bone stock, and is inserted, according to methods known in the art for insertion of PLIF implants. Preferably, the device as used for PLIF applications has the following dimensions similar to the following, see side view figure 8D: a width **811** of approximately 7 to 12 mm, and preferably about 9.4 to about 10 mm; a top dimension **812** of about 4 to 5 mm; a bottom dimension **813** of about 4-6 mm and preferably about 5 mm; a flat surface of **814** of about 4-7 mm, and preferably about 5.5 mm; a width **815** of about 5-7 mm and preferably about 5 mm; a curvature that defines an angle **816** of between about 60 and 75 degrees, and preferably about 67 degrees. See figure 8E, rear view: a length **L** of about 20 to 26 mm; a height **H** of between about 7.5 and 14.5 mm; preferably, heights of about 8, 10, 12, and 14 mm are produced with lengths of about 20 and 26 mm; desirably, the side faces **817** are machined to display a rough, ridged or grooved surface so that when the anterior end **818** of the PLIF implant is properly seated in place, ridges directed to the posterior end **819** of the PLIF implant prevent backing out of the implant. A detail of one embodiment of such a ridged surface is shown in figure 8F, wherein the following dimensions are preferred: an angle **820** for each tooth of between about 30 and 40 degrees, preferably about 35 degrees; a distance between tooth crests **821** of about 1-2 mm, preferably about 1.5 mm; a tooth crest width **822** of about 0.1 to about 0.2 mm, preferably about 0.125 mm; and a tooth height **823** of between about 0.1 to about 1 mm and preferably about

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0.5 mm; returning to figure 8E, the implant preferably has an anterior end width **824** of about 7-13 mm, preferably about 9-13 mm, with a taper angle **825** from the height **H** of about 30 to 40 degrees, preferably about 35 degrees; an instrument attachment means, **826**, such as a tapped instrument attachment hole, is provided in the posterior face of the PLIF implant; this feature is best seen from the posterior view 8G, which shows: an instrument attachment hole **826** having a diameter of about 1.5 to about 2.5 mm, preferably about 2 mm, and a depth of about 4-5 mm, preferably about 4.5 mm; an edge to center of the instrument attachment hole dimension **827** is carefully defined to match dimensions of any implant insertion device used in combination with this embodiment of the PLIF implant; a center of the instrument attachment hole to edge dimension **828** is about 4-6 mm, preferably about 5 mm, with a ridge **829** of about 1 mm running along three edges of the posterior face of the implant. In displaying the section A-A from figure 8D in figure 8E, a slight air gap **830** is shown as the section would exit bone on the concave surface of the implant and then reenter bone.

In use, the implant **810** is inserted on either side of lumbar intervertebral spaces to thereby stabilize and assist in fusion of adjacent lumbar vertebrae. This is accomplished by distraction of the lumbar vertebrae, removal of an appropriate amount and shape of intervertebral disc matter, and insertion of the implant **810**, preferably on each side on a posterior approach, according to methods known in the art. The concave surface of each implant **810** is set to face inwardly, toward the center of the vertebral body, while the convex surface of the implant **810** is set to match, as much as possible, the natural external curvature of the lumbar vertebrae.

In an analogous but alternate method for production of the cervical implant, unitary implants may be produced by appropriately sectioning and machining the anterior margin of the tibia or linea aspera of the femur. Thus, as shown in figure 10A, a left femur **1000** (posterior aspect), or in figure 10B, a left tibia **1001** (anterior aspect), is sectioned at **1004** and **1005** to remove the head, neck and greater trochanter **1002** and internal and internal condyles **1006** of the femur, or tubercle and tuberosity **1003** malleolus **1007** of the tibia. The result from such sectioning is the production of a shaft, or diaphysis, of the femur **1008** or tibia **1009**. Further processing according to this aspect of the invention involves the line asper **1010** of the femur or the anterior margin of the tibia **1011**, as shown in figure 10C. Whether produced from the femur or tibia, a diaphysial shaft **1012**, extending as shown at **1016** to a length permitted by the length of the shaft produced by the sectioning at **1004/1005**. The shaft comprises the natural intramedullary canal **1013**. The thus produced shaft is

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then further sectioned in a plane shown at **1014** to produce a shaft of bone removed from the natural intramedullary canal **1013** having a cylindrical but somewhat triangular external shape. Into this shaft may be drilled a cannulation **1015**, as shown in figure 11.

Figure 11 shows the substantially triangular shaft of substantially cortical bone **1017** produced by sectioning the shaft of the long bone down the plane **1014**. Into the shaft of bone **1017** may be drilled a bore to produce a cannulation **1015** of appropriate dimensions. The cannulation **1015** may be introduced into the unitary shaft of bone **1017** or it may be introduced into sub-segments thereof by first cutting the shaft **1017** at **1035**. In either case, the diameter of the cannulation **1015** should be limited such that at the narrowest portion between the cannulation and the wall of the device **1020** never falls below about 2 mm. When sectioned at **1035**, for example by mounting the shaft **1017** on a lathe and contacting the spinning shaft with a very narrow blade (i.e. a parting tool of about 1 mm or less width), implant blanks **1030** and **1040** are produced which may be further machined to achieve desired external and internal profiles and key way features, as described for the implant of this invention produced by alternate methods described hereinabove. Implants of any desired height, for example 5 mm to about 14 mm, may thus be produced. Figures 12-17 show specific embodiments of the implant of this invention produced according to this aspect of the invention.

Per figures 12-17, there is provided views of five different cortical bone implants according to this invention having particular geometries by way of example and not limitation. In each figure, view A is a top view, view B is a side view, view C is a detail of the grooves which angle toward the posterior aspect of the implant, and view D is a sectional view through the line A-A shown in view A. In addition, where an osteogenic plug, such as a cancellous plug is present, this is shown in view E as a top view and view F as a side view of the cancellous plug. In figure 12, there is shown an example of an implant having a height **H1** between about 5mm and about 9mm, a width **W1** of about 11 mm, and a width **W2** of about 11 mm. An outer dotted profile provides a means for comparing the shape of the implant produced according to this alternate manufacturing method with the external profile of the implant of figure 6. As can be seen, the implant produced according to this aspect of the invention has a substantially diamond-shaped external profile, as a result of the geometry of the starting shaft **1017** of bone stock.

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Figure 13 shows a device similar to that of figure 12, with a cancellous plug inserted therein. Figure 14 shows a device having a width **W1** of about 14 mm and a height **H1** of between about 5 mm and about 14 mm. Figure 15 shows a device similar to that of figure 14 with a cancellous plug inserted therein. Figure 16 shows a device having a width **W1** of about 14 mm, a width **W2** of about 14 mm, and a height of between about 5 mm and 11 mm. Figure 17 shows a device similar to that of figure 16 having a cancellous plug inserted therein. Table I below summarizes the various features and provides examples of specific dimensions for various embodiments of the implant of this invention shown in figures 12-17: Table I:

Code	Description	Figure #					
		12	13	14	15	16	17
D1	Inner Hole Diameter	5.5	5.5	5.5	5.5	5.5	5.5
D2	Hole Centerline Distance	5.5	5.5	5.5	5.5	5.5	5.5
D3	Key Way Centerline Distance	2.9	2.9	2.9	2.9	4.4	4.4
G1	Groove Depth	0.5	0.5	0.5	0.5	0.5	0.5
G2	Top Flat Width	0.5	0.5	0.5	0.5	0.5	0.5
G3	Groove Pitch width without G5	1.375	1.375	1.375	1.375	1.375	1.375
G4	Groove Pitch width with G5	1.5	1.5	1.5	1.5	1.5	1.5
G5	Flat Width	0.125	0.125	0.125	0.125	0.125	0.125
G6	Groove Angle (degrees)	30	30	30	30	30	30
H1	Overall Height	5-9	5-9	5-13	5-13	5-11	5-11
R1	Keyway Radius	0.8	0.8	0.8	0.8	0.8	0.8
R2	Corner Radius	1.5	1.5	1.5	1.5	1.5	1.5
R3	Outside Arch Radius	11	11	11	11	14	14
W1	Outside Width	11	11	14	14	14	14
W2	Outside Width	11	11	11	11	14	14
W3	Wall Thickness	≥2	≥2	≥2	≥2	≥2	≥2
CP1	Cancellous Plug Height	N/A	5-9	N/A	5-13	N/A	5-11
CP2	Cancellous Plug Diameter	N/A	5.7-5.8	N/A	5.7-5.8	N/A	5.7-5.8

4.4 Manner of Using the Implant

In use, the implant 100 is inserted into a space formed between adjacent vertebrae that are required to be fused. This may be accomplished by the surgeon removing portions of the intervertebral disk, (partial discectomy) and retracting the adjacent vertebrae to allow insertion of an appropriately dimensioned implant. The rear end 105 of the implant is inserted first, and where

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present, the external feature 120 prevents backing out of the implant. Where no external feature 120 has been inscribed into the top and bottom surfaces of the implant, it may be necessary to affix the implant in position with plate and screw retention systems known in the art. According to this invention, implants are provided having a height of between about 7 and 14 mm, a length of
5 between about 11 and 14 mm and a width of between about 11 and 14 mm. Any permutation or combination of these dimensions may be envisioned, for example (in order of height, length, width): 7x11x11, 8x11x11, etc.; 7x14x14, 8x14x14, etc.; 7x11x14, 8x11x14, etc.

Preferably, the surgeon performing the implantation saves the autologous material and debris produced in the course of the partial discectomy for packing into the canal of the present implant. In
10 addition, or alternatively, the canal may be packed (either during the surgical procedure or the canal may be pre-packed) with osteogenic, osteoinductive, or osteoconductive materials, including but not limited to: allograft bone, autograft bone, autogenous osteogenic materials including bone marrow cancellous bone and the like, demineralized bone, freeze-dried demineralized bone, Grafton (demineralized bone in glycerol), bone powder, bone derivatives, bone morphogenetic protein
15 (purified or recombinant), antibiotic, bioactive glass, hydroxyapatite, bioactive ceramics, or combinations thereof.

Following implantation, the recipient (whether human or animal) is monitored for implant stability and success in fusion. Fusion is achieved over the course of several weeks to several months, during which time increasing levels of load may be placed on the spine.

5.0 References

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U.S. Patent No. 5,306,309

6.0 What is Claimed is:

1. A spinal spacer for engagement between vertebrae, comprising:
a body formed of a bone composition and including a first end, an opposite second end, a superior face defining a superior vertebral engaging surface and an inferior face defining an inferior vertebral engaging surface; and
at least one of said vertebral engaging surfaces defining a first set of migration resistance grooves, each of said grooves including a first face defining an angle of no more than about 90 degrees relative to said one of said engaging surfaces and a second opposing sloped face, said first and second faces defining a substantially arcuate pocket therebetween for trapping vertebral bone.
2. The spacer of claim 1 wherein said grooves of said first set are arranged in series.
3. The spacer of claim 2 wherein each of said sloped faces is sloped toward said first end.
4. The spacer of claim 1 wherein said at least one of said vertebral engaging surfaces defines a peak between each of said grooves, said peak defining a flattened surface.
5. The spacer of claim 3 further comprising a second set of migration resistance grooves defined in series in at least one of said vertebral engaging surfaces, each of said grooves of said second set including a first face defining an angle of no more than about 90 degrees relative to said one of said engaging surfaces and a second opposing sloped face, said first and second faces of each of said grooves of said second set defining a pocket therebetween for trapping vertebral bone, each of said sloped faces of said second set sloping towards said second end.
6. The spacer of claim 5 wherein each of said grooves has a depth below said at least one of said vertebral engaging surfaces and said grooves of said first set are deeper than said grooves of said second set.

1 7. The spacer of claim 5 wherein each of said grooves has a depth below said at least one of
2 said vertebral engaging surfaces and said grooves of said second set are deeper than said grooves of
3 said first set.

4
1 8. The spacer of claim 1 wherein the pocket defined by said first and second faces is circularly
2 rounded.

3
1 9. The spacer of claim 1 wherein said first face is perpendicular to said at least one of said
2 vertebral engaging surfaces.

1
1 10. The spacer of claim 1 further comprising a cutting edge between said first face and said at
2 least one of said vertebral engaging surfaces.

1
1 11. The spacer of claim 4 wherein said first face at said pocket has a first height which is taller
2 than a second height of said second face at said pocket, and said peak is sloped.

3
1 12. The spacer of claim 1 wherein said superior face defines a first opening and said inferior face
2 defines a second opening, each of said openings in communication with a chamber formed through
3 said body.

4
1 13. The spacer of claim 12 wherein said first end defines a convexly curved surface.

2
1 14. The spacer of claim 13 wherein said second end is flat.

2
1 15. A hollow spinal spacer for engagement between vertebrae, comprising:
2 a body formed of bone composition and including an anterior wall with opposite ends and defining a
3 convexly curved anterior surface, an opposite posterior wall having opposite ends and defining a flat
4 posterior surface, two lateral walls, each integrally connected between said opposite ends of said
5 anterior and posterior walls to define a chamber, said walls further defining a superior vertebral
6 engaging surface defining a first opening, said first opening in communication with said chamber,

7 and an inferior vertebral engaging surface defining a second opening, said second opening in
8 communication with said chamber; and
9 at least one of said vertebral engaging faces defining a set of migration resistance grooves, each of
10 said grooves including a first face defining an angle of no more than about 90 degrees relative to
11 said one of said engaging surfaces and a second opposing sloped face, said first and second faces
12 defining a substantially arcuate pocket therebetween for trapping vertebral bone, said grooves in
13 series with said sloped faces sloping towards said anterior wall.

14
1 16. A spinal spacer for engagement between vertebrae, comprising:

1 a body formed of a bone composition and including a first end, an opposite second end, a superior
2 face defining a superior vertebral engaging surface and an inferior face defining an inferior vertebral
3 engaging surface; and
4 at least one of said vertebral engaging surfaces defining a first set of migration resistance grooves,
5 each of said grooves including a first face defining an angle of no more than about 90 degrees
6 relative to said one of said engaging surfaces and a second opposing sloped face, said first and
7 second faces defining a pocket therebetween for trapping vertebral bone, said one of said engaging
8 surfaces defining a peak between each of said grooves, said peak defining a flattened surface.

9
1 17. The spacer of claim 16 further comprising a cutting edge between said first face and said at
2 least one of said vertebral engaging surfaces.

3
1 18. The spacer of claim 16 wherein said first face at said pocket has a first height which is taller
2 than a second height of said second face at said pocket, and said peak is sloped.

3
1 19. The spacer of claim 16 wherein said first face is perpendicular to said at least one of said
2 vertebral engaging surfaces.

3
1 20. A spinal spacer for engagement between vertebrae, comprising:
2 a body formed of a bone composition and including a first end, an opposite second end, a superior
3 face defining a superior vertebral engaging surface and an inferior face defining an inferior vertebral

engaging surface; and
at least one of said vertebral engaging surfaces defining a first set of migration resistance grooves,
said at least one of said vertebral engaging surfaces defining a second set of migration resistance
grooves, each of said grooves including a first face defining an angle of no more than about 90
degrees relative to said one of said engaging surfaces and a second opposing sloped face, said first
and second faces defining a pocket therebetween for trapping vertebral bone, each of said sloped
faces of said grooves of said first set sloping toward said first end, each of said sloped faces of said
grooves of said second set sloping toward said second end.

21. The spacer of claim 20 wherein each of said grooves has a depth below said at least one of
said vertebral engaging surfaces and said grooves of said first set are deeper than said grooves of
said second set.

22. The spacer of claim 20 wherein each of said grooves has a depth below said at least one of
said vertebral engaging surfaces and said grooves of said second set are deeper than said grooves of
said first set.

23. The spacer of claim 20 wherein said at least one of said vertebral engaging surfaces defines a
peak between each of said grooves, said peak defining a flattened surface.

24. The spacer of claim 23 wherein said first face at said pocket has a first height which is taller
than a second height of said second face at said pocket, and said peak is sloped.

25. A spinal spacer consisting substantially of cortical bone, a top face, a bottom face, a canal
surrounded by a continuous or discontinuous wall of cortical bone comprising a canal surrounded by
convexly curved anterior cortical bone face and three substantially rectilinear cortical bone faces
unitary with said convexly curved anterior cortical bone face.

26. The implant of claim 25 which has a substantially "D"-shaped external profile.

1 27. The implant of claim 26 wherein said canal has a substantially "D"-shape.

1 28. The implant of claim 25 further having an external feature on said top face, said bottom face
2 or both.

1 29. A method of making at least one implant consisting substantially of cortical bone, said
2 implant comprising a canal surrounded by a continuous or discontinuous wall of cortical bone
3 having a top face and a bottom face, each of which is substantially planar, with said planes being
4 substantially parallel to each other, said method comprising:

- 5 (a) obtaining a plug of bone consisting substantially of cortical bone by using a core cutter
6 having a central drill bit, thereby forming a canal through the bone plug obtained with
7 the core cutter;
8 (b) machining the bone plug of step (a) to produce a "washer-shaped" bone plug;
9 (c) machining the canal through the bone plug to form an asymmetric shape therein; and
10 (d) using said asymmetric shape to machine an outside profile of the bone plug.

1 30. A method of making at least one implant consisting substantially of cortical bone, said
2 implant comprising a canal surrounded by a continuous or discontinuous wall of cortical bone
3 having a top face and a bottom face, each of which is substantially planar, with said planes being
4 substantially parallel to each other, said method comprising:

- 5 (a) cutting a segment of cortical bone;
6 (b) shaping said segment of cortical bone into a symmetric half of the final shape of said
7 implant comprising a canal surrounded by a continuous or discontinuous wall of cortical
8 bone, such that when implanted in juxtaposition with a mirror image segment, an implant
9 is formed having a circular, an elliptical, or an asymmetric shape, a top face and a bottom
10 face, each of which is substantially planar, with said planes being substantially parallel to
11 each other; and
12 (c) cutting appropriate lengths of said shaped segment of cortical bone such that said cut
13 length provides half of an implant having a desired height.

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1 31. A method for inducing fusion of cervical vertebrae which comprises removing a portion of
2 the intervertebral disc between the adjacent vertebrae that are to be fused, and inserting into said
3 space at least one implant consisting substantially of cortical bone, said implant comprising a canal
4 surrounded by a continuous or discontinuous wall of cortical bone having a top face and a bottom
5 face, each of which is substantially planar, with said planes being substantially parallel to each
6 other.

1 32. An implant consisting substantially of cortical bone, said implant having been prepared by a
2 process comprising:

- 3 (a) obtaining a plug of bone consisting substantially of cortical bone by using a core cutter
4 having a central drill bit, thereby forming a canal through the bone plug obtained with
5 the core cutter;
6 (b) machining the bone plug of step (a) to produce a "washer-shaped" bone plug;
7 (c) machining the canal through the bone plug to form an asymmetric shape therein; and
8 (d) using said asymmetric shape to machine an outside profile of the bone plug.
- 1

1

7.0 Abstract of the Disclosure

5 An implant composed substantially of cortical bone is provided for use in cervical Smith-Robinson vertebral fusion procedures. The implant is derived from allograft or autograft cortical bone sources, is machined to form a symmetrically or asymmetrically shaped (e.g. a substantially “D”-shaped) implant having a canal running therethrough according to methods of this invention, and inserted into the space between adjacent cervical vertebrae to provide support and induce fusion of the adjacent vertebrae. Osteogenic, osteoinductive or osteoconductive materials may be packed into the canal of the implant to expedite vertebral fusion and to allow autologous bony ingrowth.

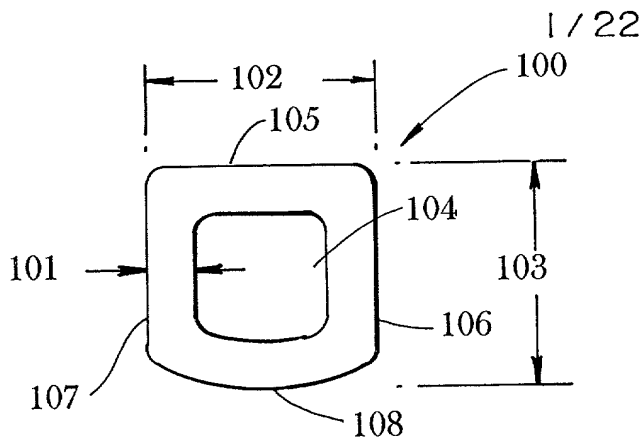


Fig. 1A

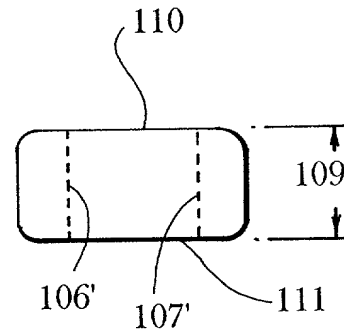


Fig. 1B

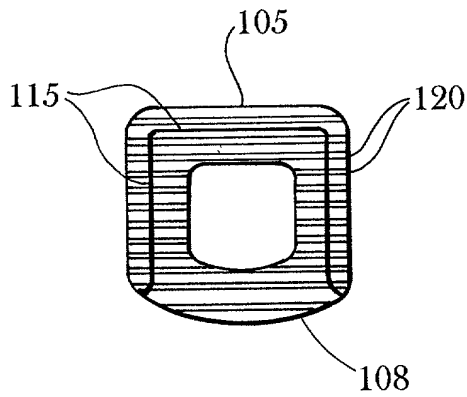


Fig. 1C

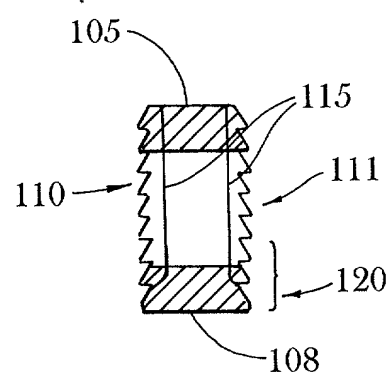


Fig. 1D

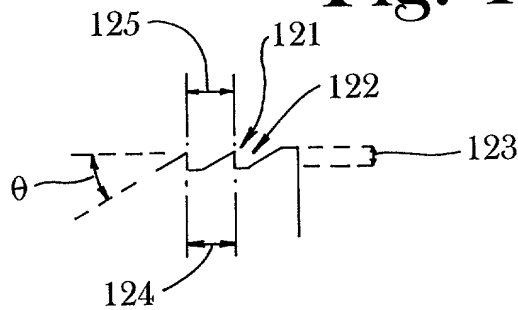


Fig. 1E

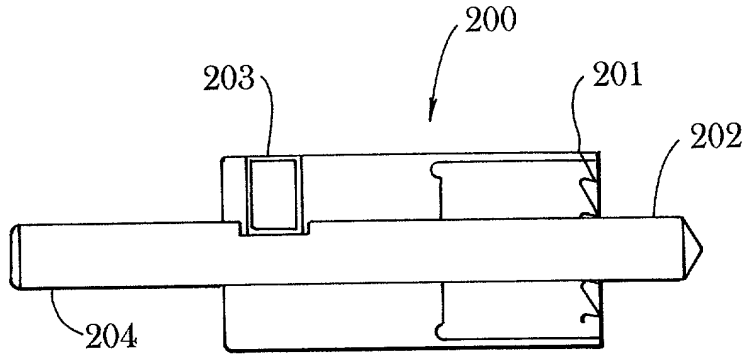


Fig. 2A

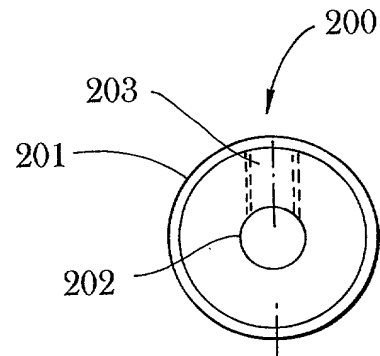


Fig. 2B

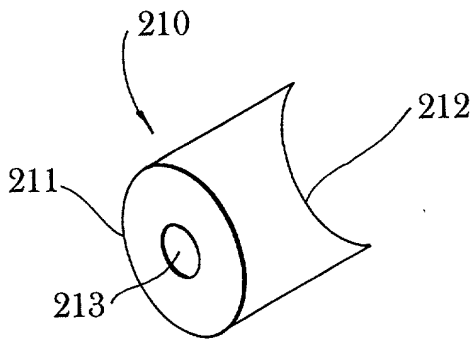


Fig. 2C

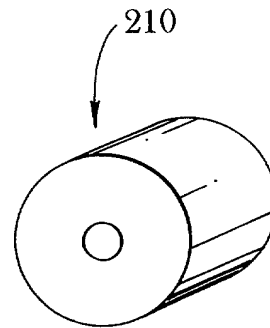


Fig. 2D

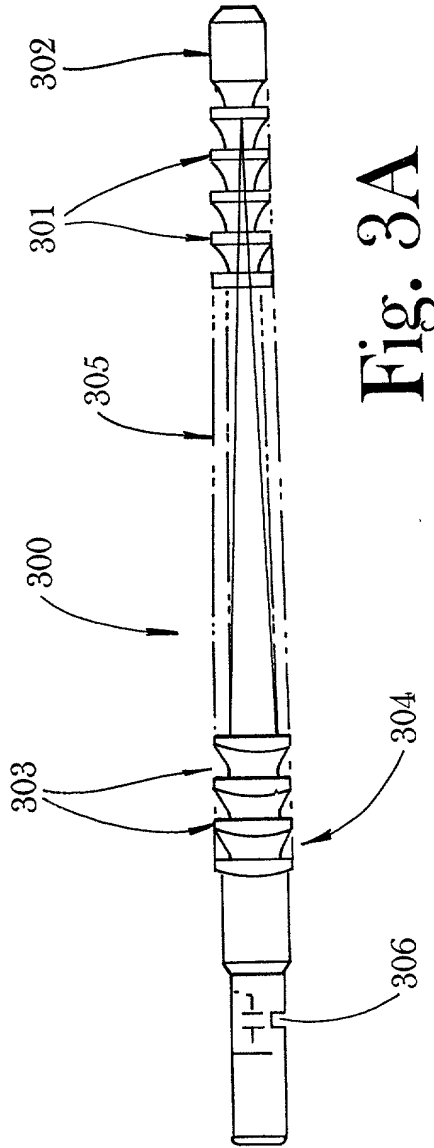


Fig. 3A

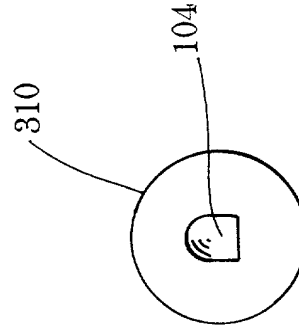


Fig. 3B

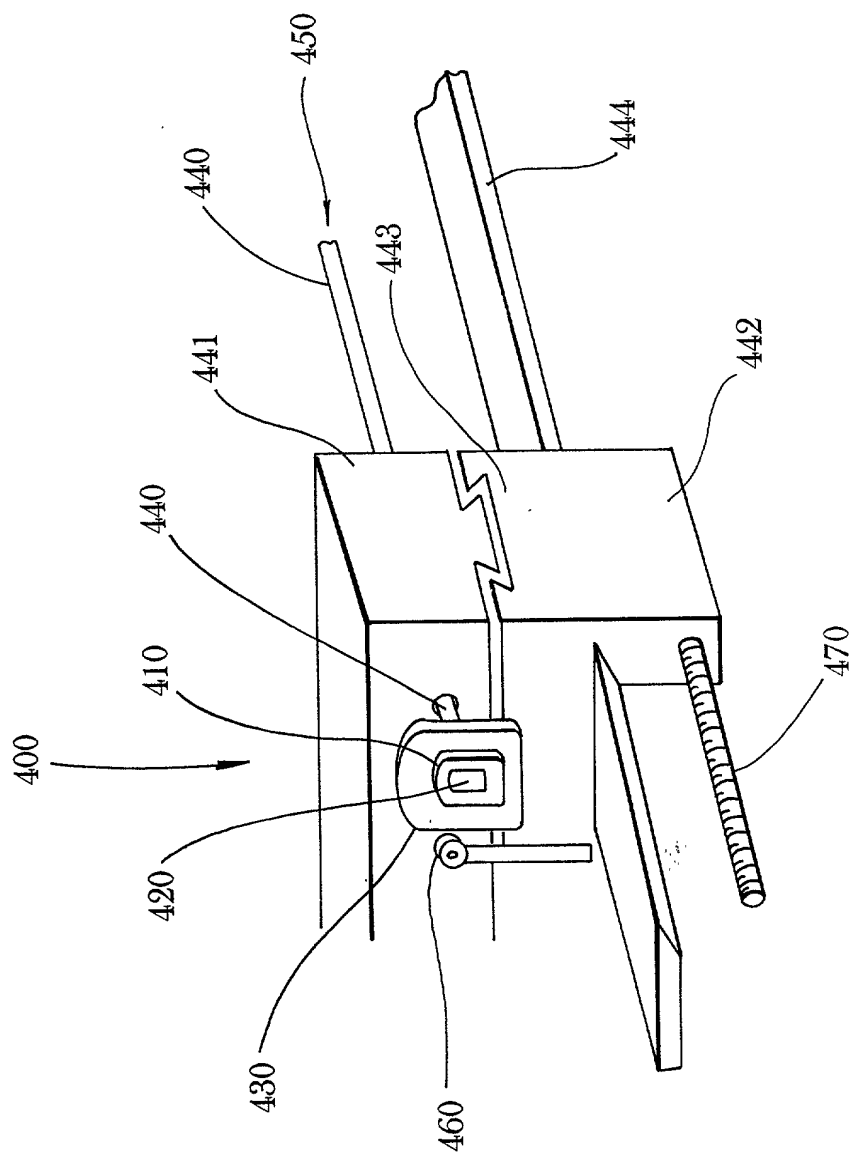


Fig. 4A

Fig. 5A

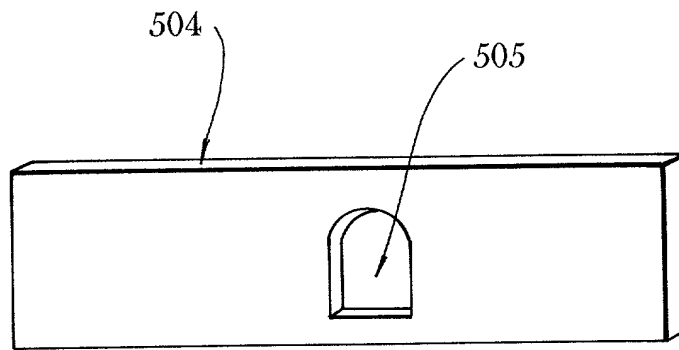
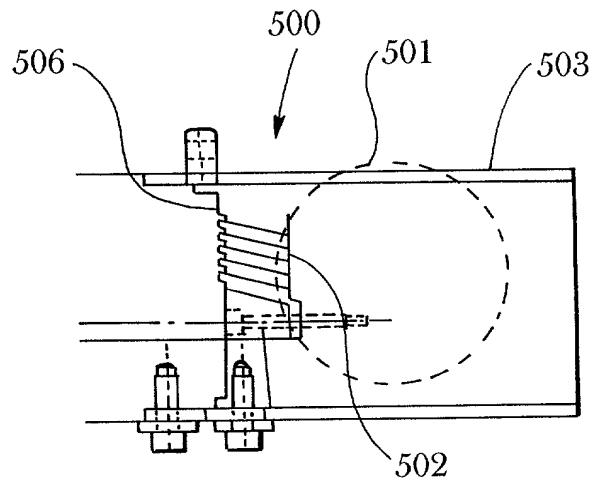


Fig. 5B

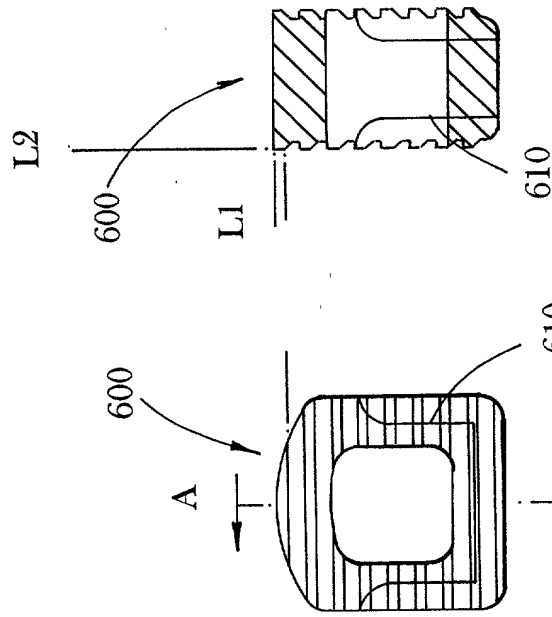


Fig. 6D

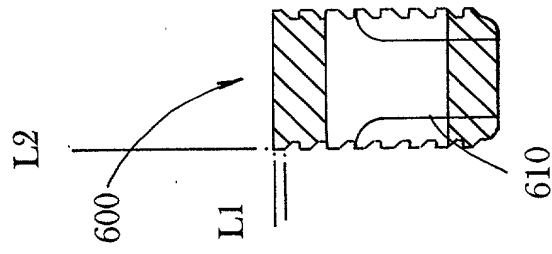


Fig. 6E

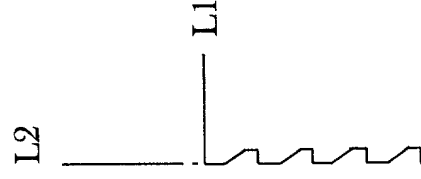


Fig. 6F

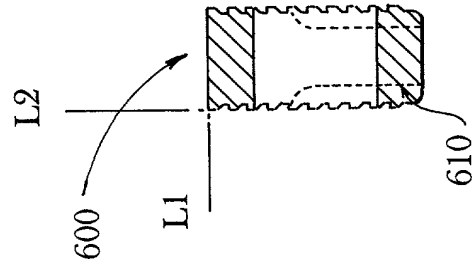


Fig. 6H

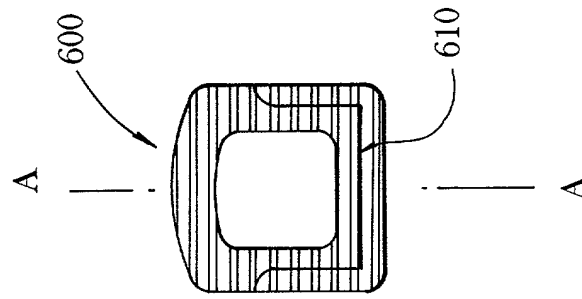


Fig. 6G

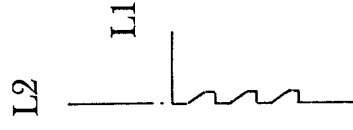


Fig. 6I

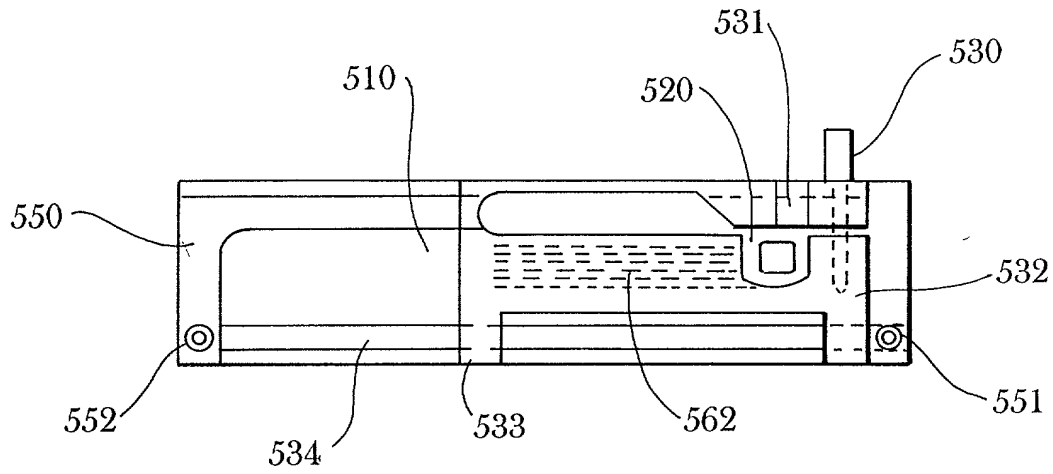


Fig. 5C

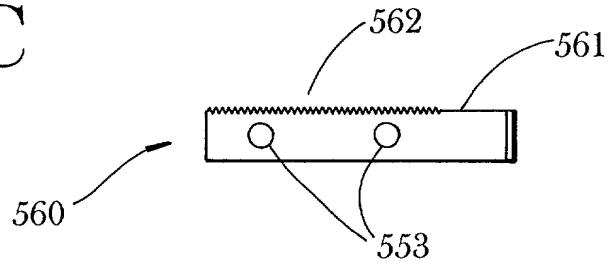


Fig. 5D

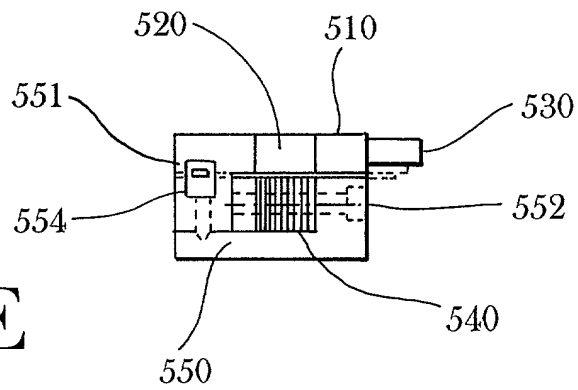


Fig. 5E

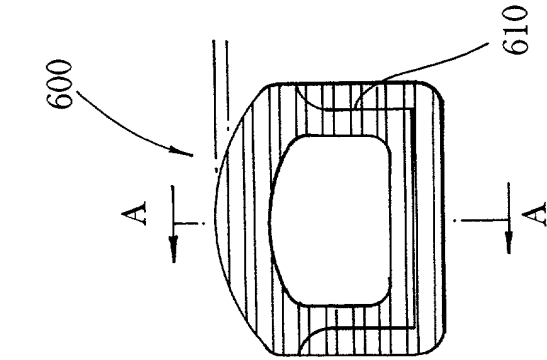


Fig. 6A

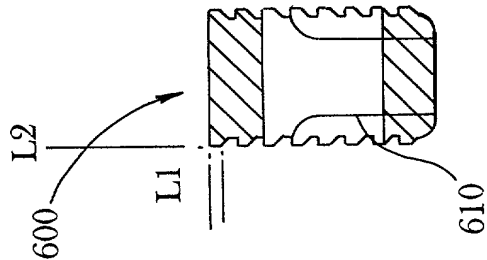


Fig. 6B

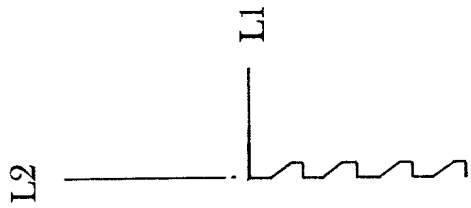


Fig. 6C

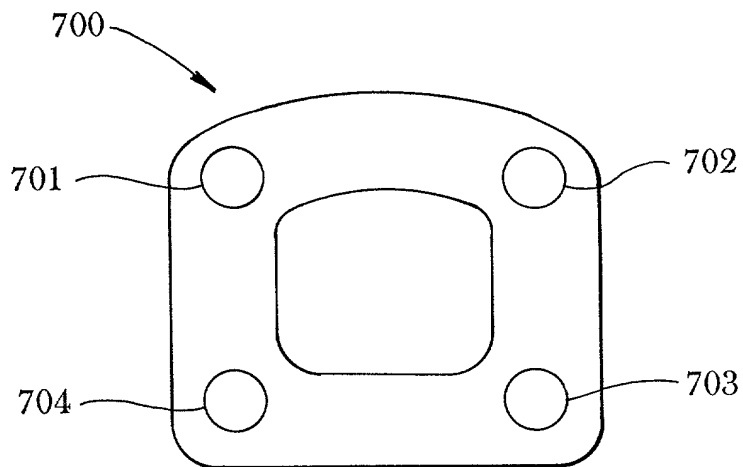


Fig. 7A

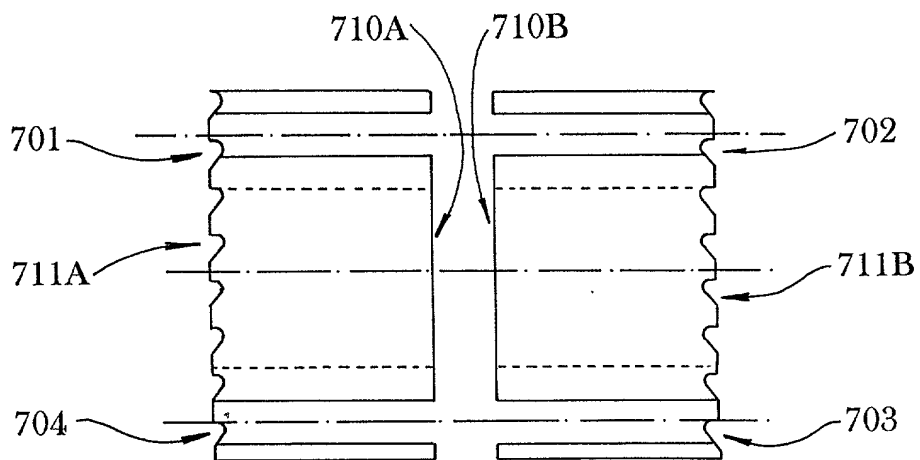


Fig. 7B

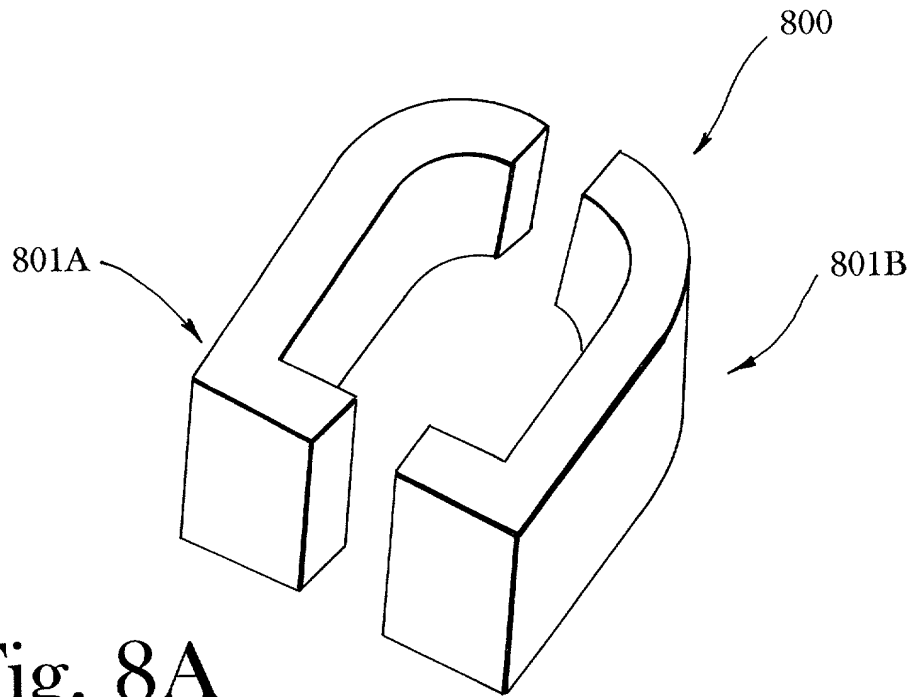


Fig. 8A

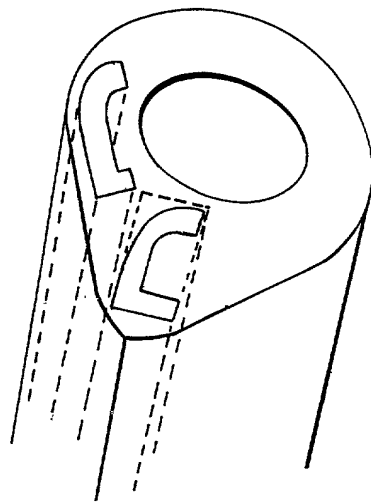


Fig. 8B

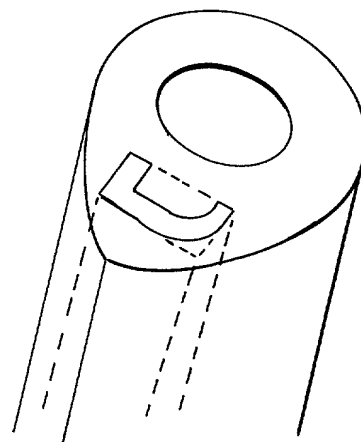


Fig. 8C

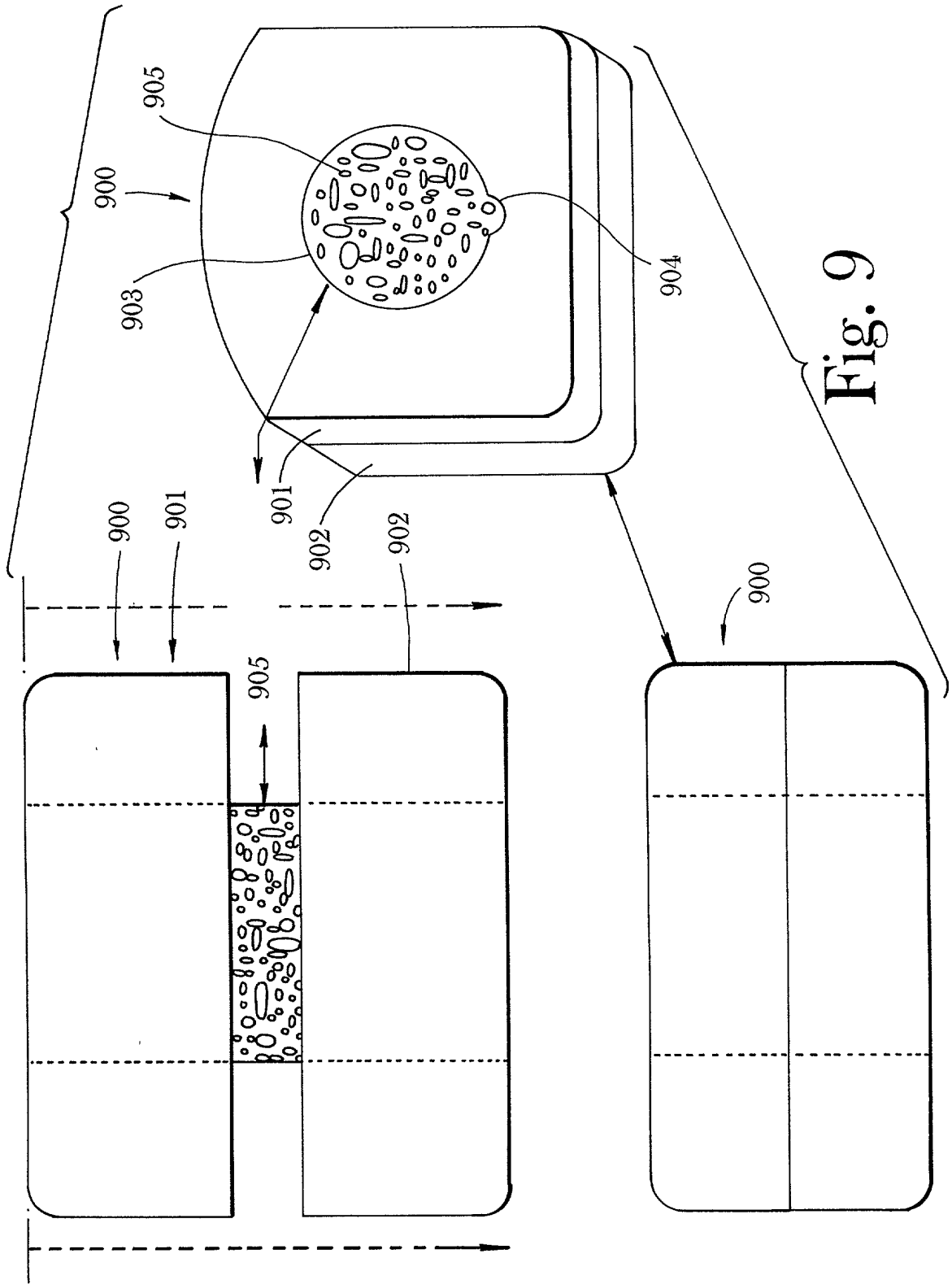


Fig. 10A

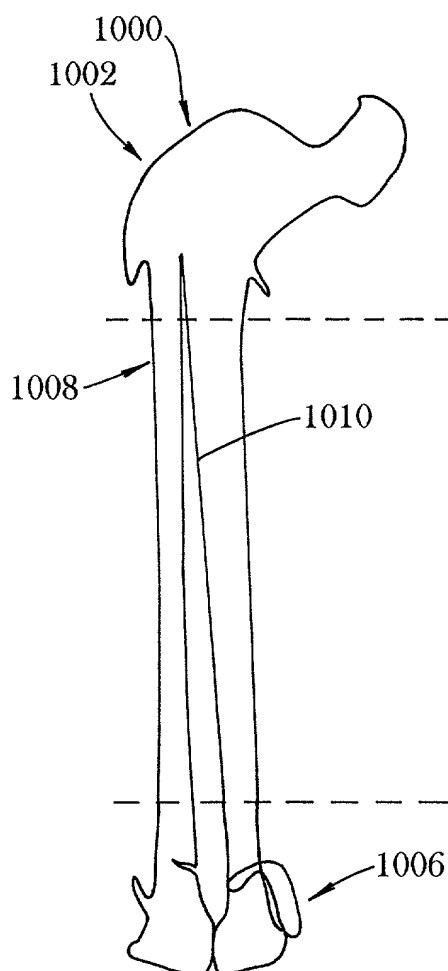


Fig. 10B

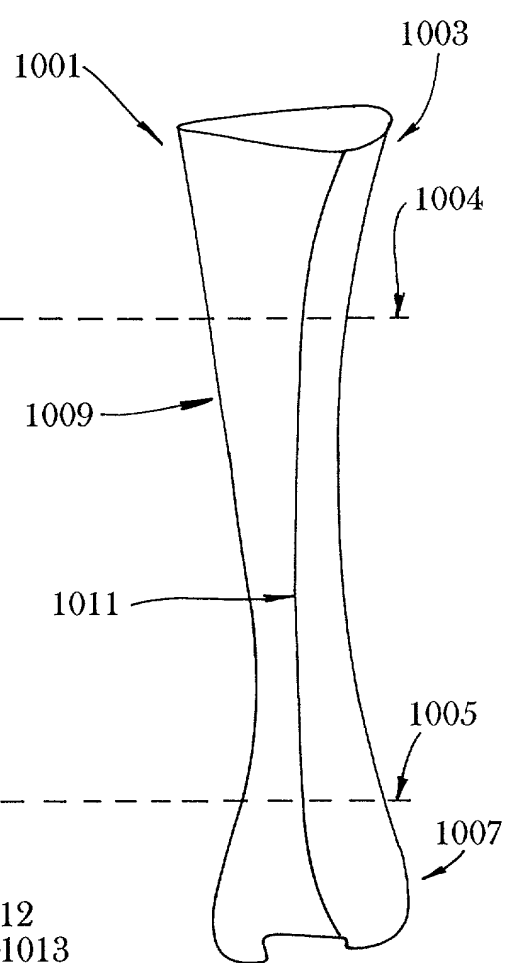
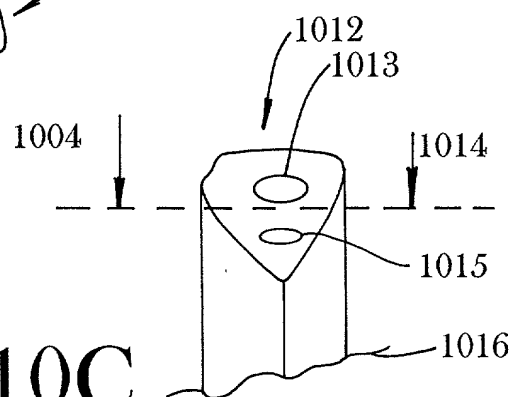


Fig. 10C



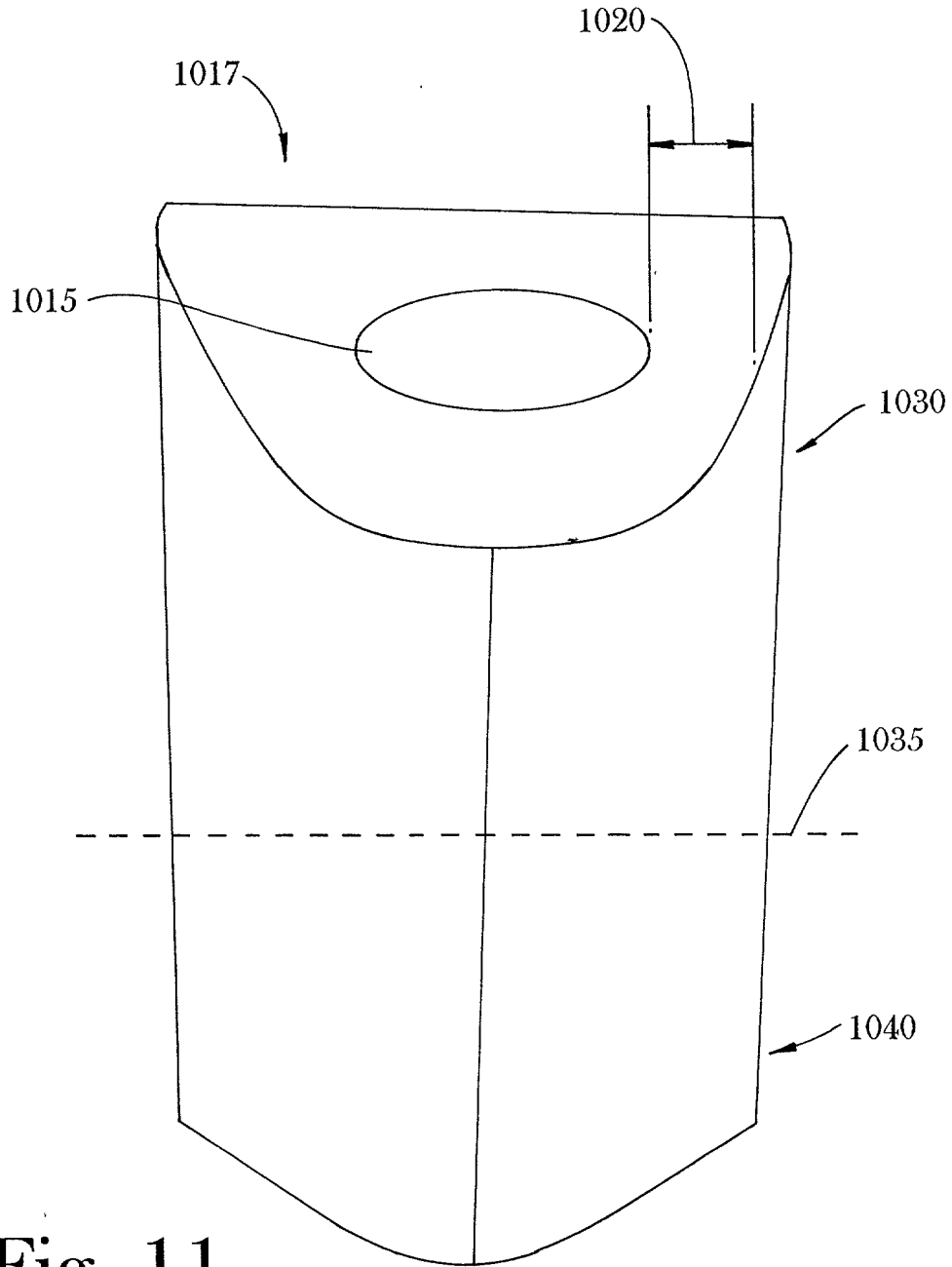


Fig. 11

Fig. 12A

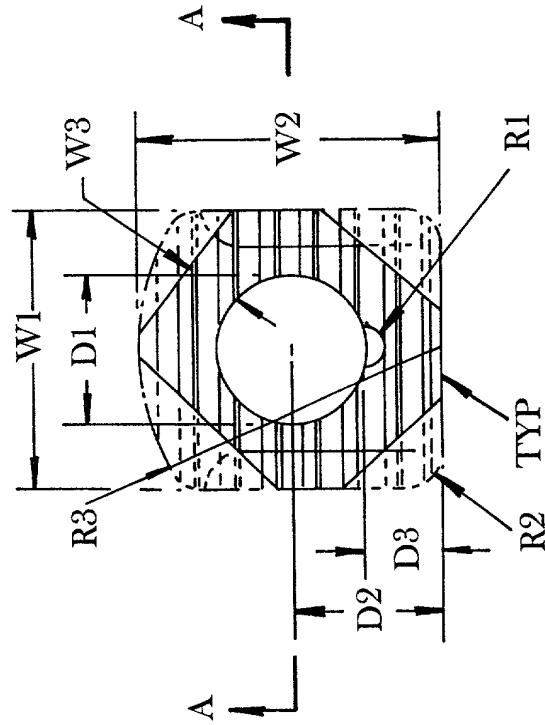


Fig. 12B

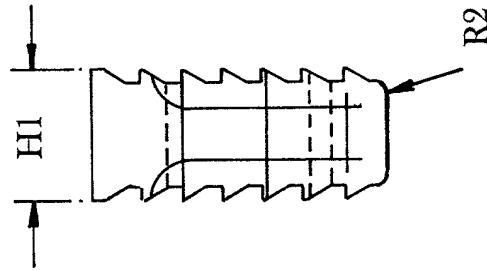


Fig. 12C

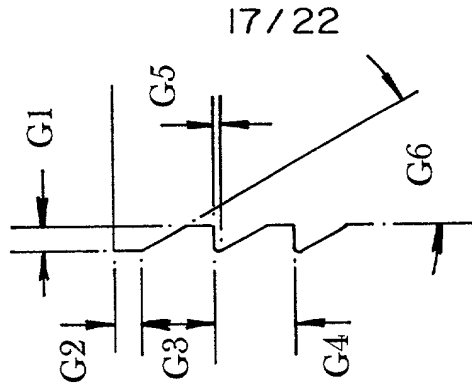
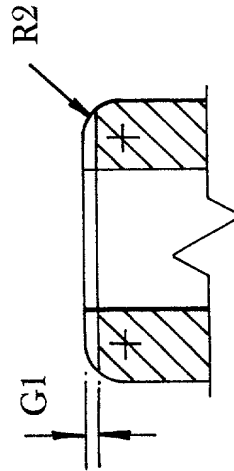


Fig. 12D



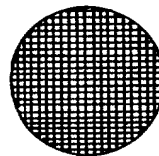
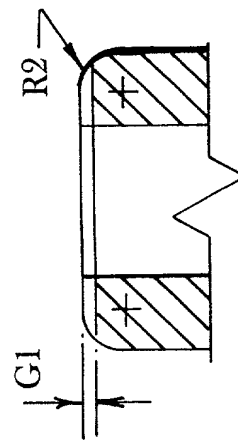
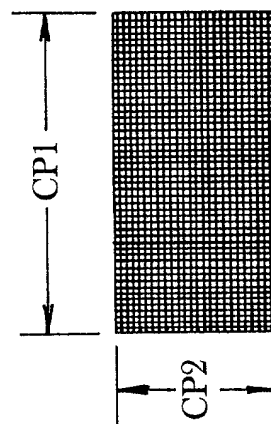
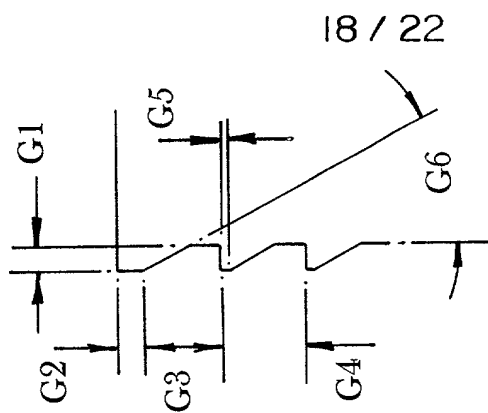
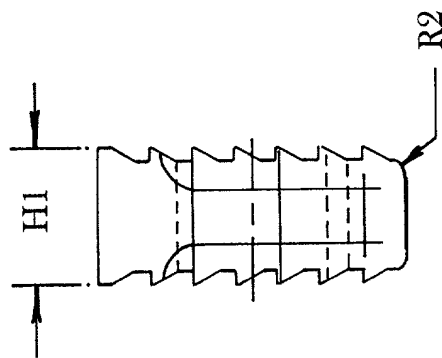
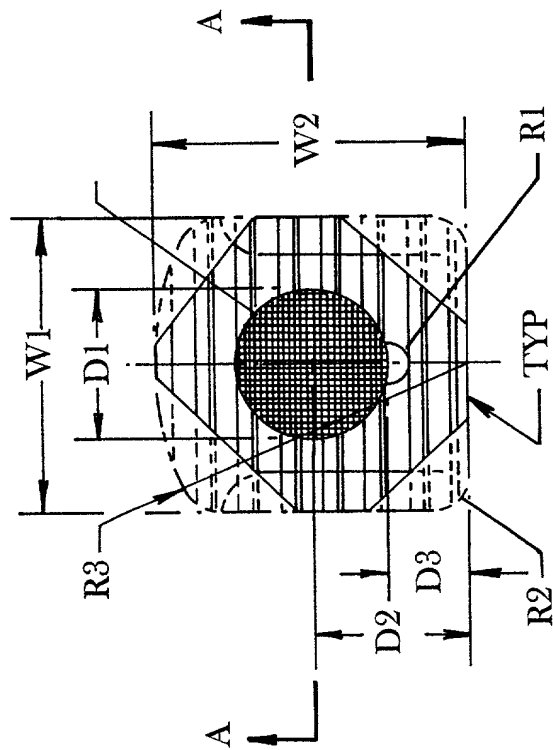


Fig. 14A

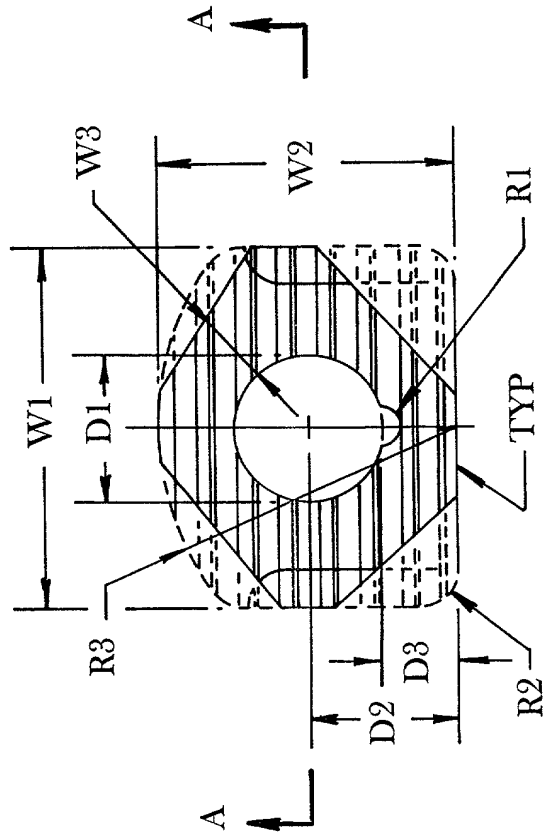


Fig. 14B

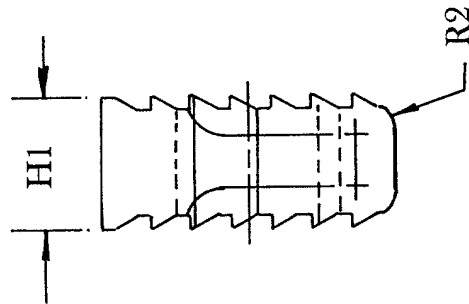


Fig. 14C

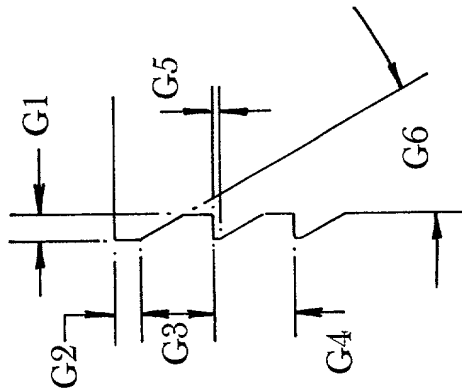


Fig. 14D

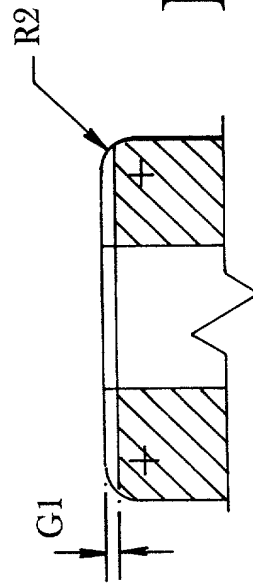


Fig. 15A

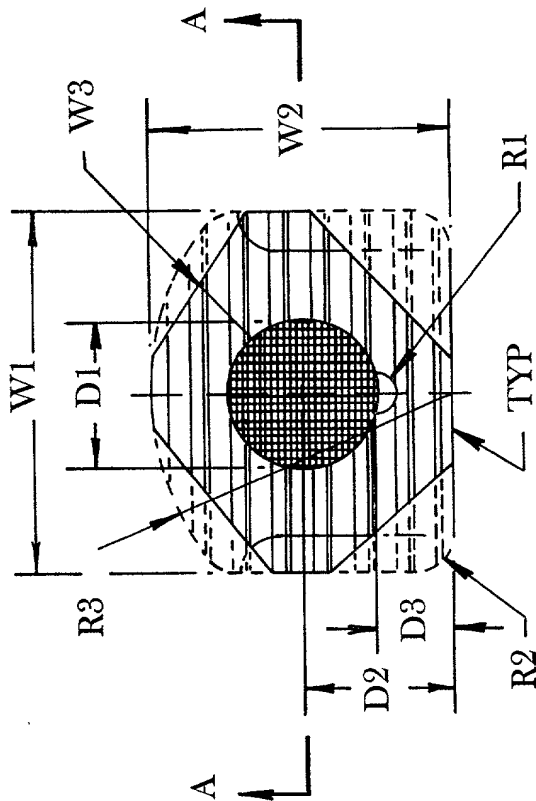


Fig. 15B

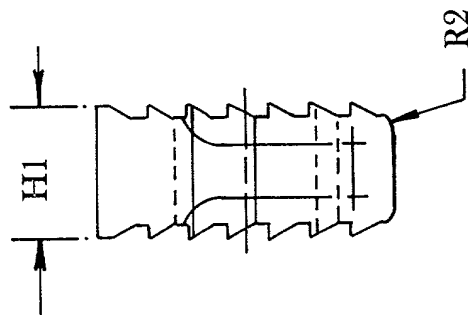


Fig. 15C

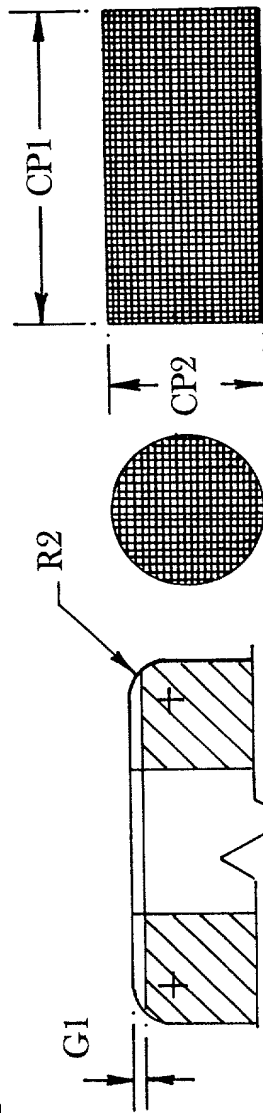
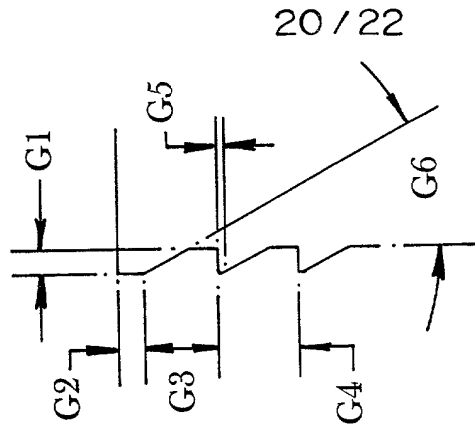


Fig. 15D

Fig. 15E

Fig. 15F

Fig. 16A

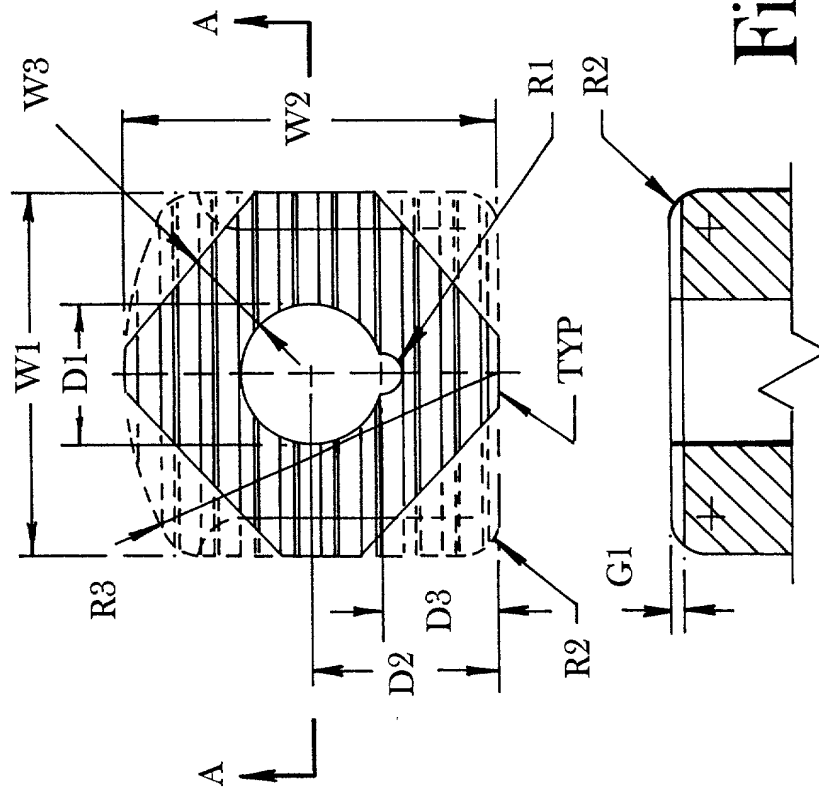


Fig. 16B

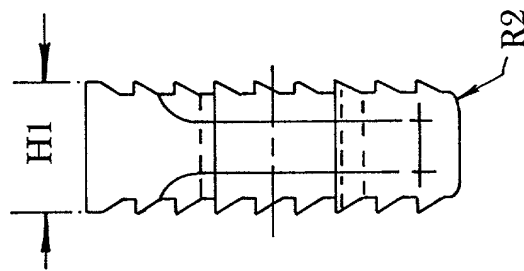


Fig. 16C

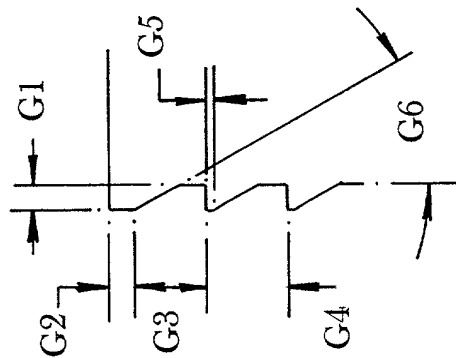


Fig. 16D

Fig. 17A

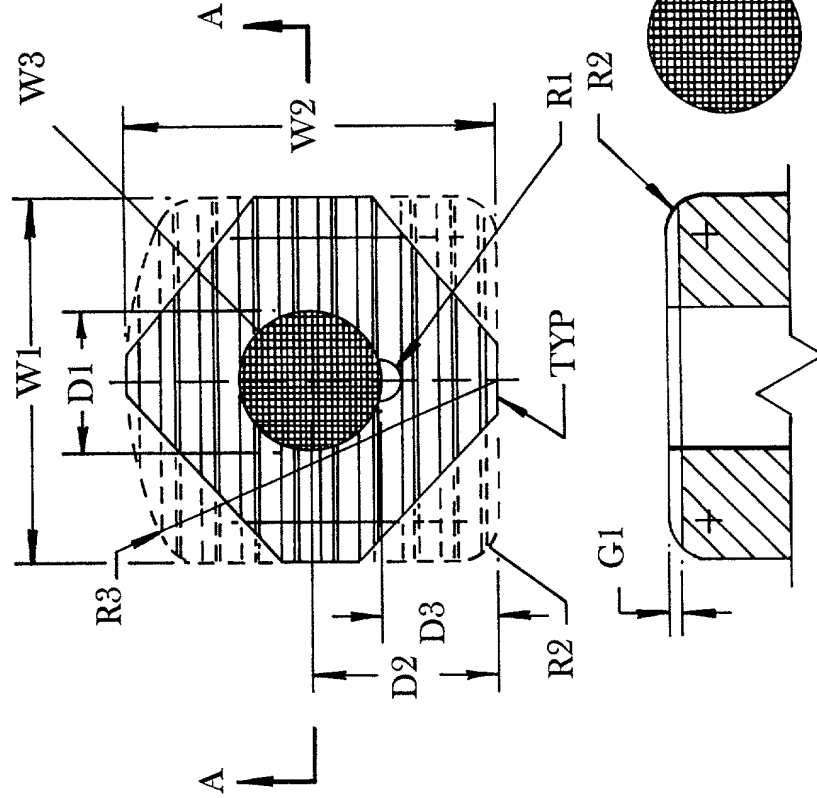


Fig. 17B

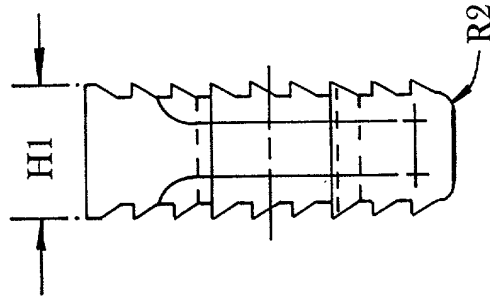


Fig. 17C

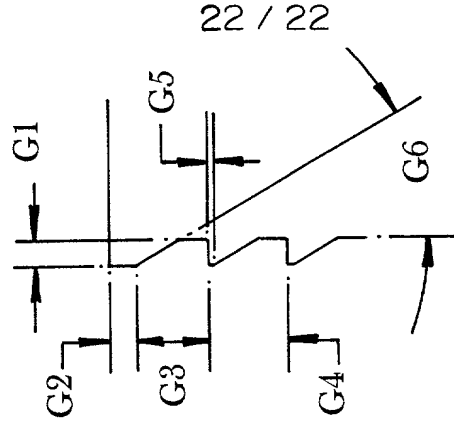


Fig. 17F

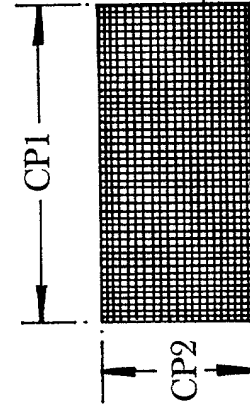
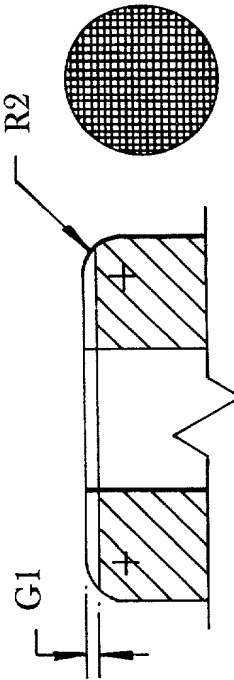


Fig. 17D Fig. 17E



PATENT APPLICATION

DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

ATTORNEY DOCKET NO. **TB-1041B**

As a below named inventor, I hereby declare that:

My residence/post office address and citizenship are as stated below next to my name;

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

Cortical Bone Cervical Smith-Robinson Fusion Implant

the specification of which is attached hereto unless the following box is checked:

☐ was filed on _____ as US Application Serial No. or PCT International Application
Number _____ and was amended on _____ (if applicable).

I hereby state that I have reviewed and understood the contents of the above-identified specification, including the claims, as amended by any amendment(s) referred to above. I acknowledge the duty to disclose all information which is material to patentability as defined in 37 CFR 1.56.

Foreign Application(s) and/or Claim of Foreign Priority

I hereby claim foreign priority benefits under Title 35, United States Code Section 119 of any foreign application(s) for patent or inventor(s) certificate listed below and have also identified below any foreign application for patent or inventor(s) certificate having a filing date before that of the application on which priority is claimed:

COUNTRY	APPLICATION NUMBER	DATE FILED	PRIORITY CLAIMED UNDER 35 U.S.C. 119
			YES: _____ NO: _____
			YES: _____ NO: _____

Provisional Application

I hereby claim the benefit under Title 35, United States Code Section 119(e) of any United States provisional application(s) listed below:

APPLICATION SERIAL NUMBER	FILING DATE

U.S. Priority Claim

I hereby claim the benefit under Title 35, United States Code, Section 120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code Section 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, Section 1.56(a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

APPLICATION SERIAL NUMBER	FILING DATE	STATUS(patented/pending/abandoned)
US98/17769	8/27/1997	Pending

POWER OF ATTORNEY:

As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) listed below to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.

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I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

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Date _____

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Date

**DECLARATION AND POWER OF ATTORNEY
FOR PATENT APPLICATION (continued)**

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Inventor's Signature _____

Date _____

Full Name of Inventor: _____

Citizenship: _____

Residence: _____

Post Office Address: _____

Inventor's Signature _____

Date _____

Full Name of Inventor: _____

Citizenship: _____

Residence: _____

Post Office Address: _____

Inventor's Signature _____

Date _____